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(57) Abstract

Thermostable glycosidase enzymes derived from various Thermococcus, Staphylothermus and Pyrococcus organisms is disclosed. The enzymes are produced from native or recombinant host cells and can be utilized in the food processing industry, pharmaceutical industry and in the textile industry, detergent industry and in the baking industry.

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GLYCOSIDASE ENZYMES

BACKGROUND OF THE INVENTION

1. Field of the Inventions

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This invention relates to newly identified polynucleotides, polypeptides encoded by such polynucleotides, the use of such polynucleotides and polypeptides, as well as the production and isolation of such polynucleotides and polypeptides. More particularly, the polynucleotides and polypeptides of the present invention has been putatively identified as glucosidases. α -galactosidases, β -galactosidases, β -mannosidases, β -mannases, endoglucanases, and pullalanases.

2. Description of Related Art

The glycosidic bond of \beta-galactosides can be cleaved by different classes of enzymes: (i) phospho-β-galactosidases (EC3.2.1.85) are specific for a phosphorylated substrate generated via phosphoenolpyruvate phosphotransferase system (PTS)-dependent uptake; (ii) typical β-galactosidases (EC 3.2.1.23), represented by the Escherichia coli LacZ enzyme, which are relatively specific for β -galactosides; and (iii) β -glucosidases (EC 3.2.1.21) such as the enzymes of Agrobacterium faecalis, Clostridium thermocellum, Pyrococcus furiosus or Sulfolobus solfataricus (Day, A.G. and Withers, S.G., (1986) Purification and characterization of a β-glucosidase from Alcaligenes faecalis. Can. J. Biochem, Cell. Biol. 64, 914-922; Kengen, S.W.M., et al. (1993) Eur. J. Biochem., 213, 305-312; Ait, N., Cruezet, N. and Cattaneo, J. (1982) Properties of β-glucosidase purified from Clostridium thermocellum. J. Gen. Microbiol. 128, 569-577; Grogan, D.W. (1991) Evidence that β-galactosidase of Sulfolobus solfataricus is only one of several activities of a thermostable β-D-glycodiase. Appl. Environ. Microbiol. 57, 1644-1649). Members of the latter group, although highly specific with respect to the \beta-anomeric configuration of the glycosidic linkage, often display a rather relaxed substrate specificity and hydrolyze βglucosides as well as β -fucosides and β -galactosides.

Generally, α -galactosidases are enzymes that catalyze the hydrolysis of galactose groups on a polysaccharide backbone or hydrolyze the cleavage of di- or oligosaccharides comprising galactose.

Generally, β-mannanases are enzymes that catalyze the hydrolysis of mannose groups internally on a polysaccharide backbone or hydrolyze the cleavage of di- or oligosaccaharides comprising mannose groups. β-mannosidases hydrolyze non-reducing, terminal mannose residues on a mannose-containing polysaccharide and the cleavage of di- or oligosaccaharides comprising mannose groups.

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Guar gum is a branched galactomannan polysaccharide composed of β -1,4 linked mannose backbone with α -1,6 linked galactose side chains. The enzymes required for the degradation of guar are β -mannanase, β -mannosidase and α -galactosidase. β -mannanase hydrolyses the mannose backbone internally and β -mannosidase hydrolyses non-reducing. terminal mannose residues. α -galactosidase hydrolyses α -linked galactose groups.

Galactomannan polysaccharides and the enzymes that degrade them have a variety of applications. Guar is commonly used as a thickening agent in food and is utilized in hydraulic fracturing in oil and gas recovery. Consequently, galactomannanases are industrially relevant for the degradation and modification of guar. Furthermore, a need exists for thermostable galactomannases that are active in extreme conditions associated with drilling and well stimulation.

There are other applications for these enzymes in various industries, such as in the beet sugar industry. 20-30% of the domestic U.S. sucrose consumption is sucrose from sugar beets. Raw beet sugar can contain a small amount of raffinose when the sugar beets are stored before processing and rotting begins to set in. Raffinose inhibits the crystallization of sucrose and also constitutes a hidden quantity of sucrose. Thus, there is merit to eliminating raffinose from raw beet sugar. α -Galactosidase has also been used as a digestive aid to break down raffinose, stachyose, and verbascose in such foods as beans and other gassy foods.

β-galactosidases which are active and stable at high temperatures appear to be superior enzymes for the production of lactose-free dietary milk products (Chaplin, M.F.

and Bucke, C. (1990) In: Enzyme Technology, pp. 159-160, Cambridge University Press, Cambridge, UK). Also, several studies have demonstrated the applicability of β-galactosidases to the enzymatic synthesis of oligosaccharides via transglycosylation reactions (Nilsson, K.G.I. (1988) Enzymatic synthesis of oligosaccharides. Trends Biotechnol. 6, 156-264; Cote, G.L. and Tao, B.Y. (1990) Oligosaccharide synthesis by enzymatic transglycosylation. Glycoconjugate J. 7, 145-162). Despite the commercial potential, only a few β-galactosidases of thermophiles have been characterized so far. Two genes reported are β-galactoside-cleaving enzymes of the hyperthermophilic bacterium *Thermotoga maritima*, one of the most thermophilic organotrophic eubacteria described to date (Huber, R., Langworthy, T.A., König, H., Thomm, M., Woese, C.R., Sleytr, U.B. and Stetter, K.O. (1986) *T. martima* sp. nov. represents a new genus of unique extremely thermophilic eubacteria growing up to 90°C, Arch. Microbiol. 144, 324-333) one of the most thermophilic organotrophic eubacteria described to date. The gene products have been identified as a β-galactosidase and a β-glucosidase.

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Pullulanase is well known as a debranching enzyme of pullulan and starch. The enzyme hydrolyzes α -1,6-glucosidic linkages on these polymers. Starch degradation for the production or sweeteners (glucose or maltose) is a very important industrial application of this enzyme. The degradation of starch is developed in two stages. The first stage involves the liquefaction of the substrate with α -amylase, and the second stage, or saccharification stage, is performed by β -amylase with pullalanase added as a debranching enzyme, to obtain better yields.

Endoglucanases can be used in a variety of industrial applications. For instance, the endoglucanases of the present invention can hydrolyze the internal β-1,4-glycosidic bonds in cellulose, which may be used for the conversion of plant biomass into fuels and chemicals. Endoglucanases also have applications in detergent formulations, the textile industry, in animal feed, in waste treatment, and in the fruit juice and brewing industry for the clarification and extraction of juices.

Brief Description of the Drawings

The following drawings are illustrative of embodiments of the invention and are not meant to limit the scope of the invention as encompassed by the claims.

Figures 1a-b are the full-length DNA and corresponding deduced amino acid sequence of M11TL of the present invention. Sequencing was performed using a 378 automated DNA sequencer for all sequences of the present invention (Applied Biosystems, Inc.).

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Figure 2 is an illustration of the full-length DNA and corresponding deduced amino acid sequence of OC1/4V-33B/G.

Figure 3 is an illustration of the full-length DNA and corresponding deduced amino acid sequence of F1-12G.

Figures 4a-b are the full-length DNA and corresponding deduced amino acid sequence of 9N2-31B/G.

Figures 5a-b are the full-length DNA and corresponding deduced amino acid sequence of MSB8-6G.

Figure 6 is the full-length DNA and corresponding deduced amino acid sequence of AEDII12RA-18B/G.

Figures 7a-b are the full-length DNA and corresponding deduced amino acid sequence of GC74-22G.

Figures 8a-b are the full-length DNA and corresponding deduced amino acid sequence of VC1-7G1.

Figures 9a-c are the full-length DNA and corresponding deduced amino acid sequence of 37GP1.

Figures 10a-c are the full-length DNA and corresponding deduced amino acid sequence of 6GC2.

Figures 11a-d are the full-length DNA and corresponding deduced amino acid sequence of 6GP2.

Figures 12a-c are the full-length DNA and corresponding deduced amino acid sequence of 63GB1.

Figures 13a-b are the full-length DNA and corresponding deduced amino acid sequence of OC1/4V.

Figures 14a-e are the full-length DNA and corresponding deduced amino acid sequence of 6GP3.

Figures 15a-d are the full-length DNA and corresponding deduced amino acid sequence of *Thermotoga maritima* MSB8-6GP2.

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Figures 16a-c are the full-length DNA and corresponding deduced amino acid sequence of *Thermotoga maritima* MSB8-6GB4.

Figures 17a-d are the full-length DNA and corresponding deduced amino acid sequence of *Banki gouldi* 37GP4.

Figures 18a-b are the full-length DNA and corresponding deduced amino acid-sequence of *Pyrococcus furiosus* VC1-7EG1.

SUMMARY OF THE INVENTION

In a preferred embodiment of the present invention, there are provided isolated nucleic acids (polynucleotides) which encode mature enzymes having the deduced amino acid sequences of Figures 1-18 (SEQ ID NOS: 15-28 and 61-64).

In another embodiment, the invention provides a method for producing a polypeptide including culturing host cells containing the polynucleotide of Figures 1-18 and expressing from the host cell a polypeptide encoded by the polynucleotide and isolating the polypeptide.

In another embodiment, the invention provides an enzyme selected from the group consisting of an enzyme having an amino acid sequence set forth in SEQ ID NOS: 15-28 or 61-64 and an enzyme which has at least 30 consecutive amino acid residue as an enzyme having an amino acid sequence set forth in SEQ ID NOS: 15-28 or 61-64.

In yet another embodiment, the invention provides a method for generating glucose from soluble cell oligosaccharides which includes contacting a sample containing oligosaccharides with an effective amount of an enzyme selected from the group of

enzymes having the amino acid sequence set forth in SEQ ID NOS: 15-28, 61-63 and 64 such that glucose is produced

The publications discussed herein are provided solely for their disclosure prior to the filing date of the present application. Nothing herein is to be construed as an admission that the invention is not entitled to antedate such disclosure by virtue of prior invention.

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Definitions

"Monosaccharide", as used herein, refers to a single polyhydroxy aldehyde or ketone unit.

"Oligosaccharide", as used herein, consist of short chains of monosaccharide units joined together by covalent bonds. Of these, the most abundant are the disaccharides, which have two monosaccharide units.

"Polysaccharide", as used herein, consists of long chains having many monosaccharide units.

The term "gene" means the segment of DNA involved in producing a polypeptide chain; it includes regions preceding and following the coding region (leader and trailer) as well as intervening sequences (introns) between individual coding segments (exons).

A coding sequence is "operably linked to" another coding sequence when RNA polymerase will transcribe the two coding sequences into a single mRNA, which is then translated into a single polypeptide having amino acids derived from both coding sequences. The coding sequences need not be contiguous to one another so long as the expressed sequences ultimately process to produce the desired protein.

"Recombinant" enzymes refer to enzymes produced by recombinant DNA techniques; *i.e.*, produced from cells transformed by an exogenous DNA construct encoding the desired enzyme. "Synthetic" enzymes are those prepared by chemical synthesis.

A DNA "coding sequence of" or a "nucleotide sequence encoding" a particular enzyme, is a DNA sequence which is transcribed and translated into an enzyme when placed under the control of appropriate regulatory sequences.

Detailed Description of the Invention

The polynucleotides and polypeptides of the present invention have been identified as glucosidases, α -galactosidases, β -galactosidases, β -mannosidases, β -mannanases, endoglucanases, and pullalanases as a result of their enzymatic activity.

In accordance with one aspect of the present invention, there are provided novel enzymes, as well as active fragments, analogs and derivatives thereof.

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In accordance with another aspect of the present invention, there are provided isolated nucleic acid molecules encoding the enzymes of the present invention including mRNAs, cDNAs, genomic DNAs as well as active analogs and fragments of such enzymes.

In accordance with yet a further aspect of the present invention, there is provided a process for producing such polypeptides by recombinant techniques comprising culturing recombinant prokaryotic and/or eukaryotic host cells, containing a nucleic acid sequence of the present invention, under conditions promoting expression of said enzymes and subsequent recovery of said enzymes.

In accordance with yet a further aspect of the present invention, there is provided a process for utilizing such enzymes, or polynucleotides encoding such enzymes for hydrolyzing lactose to galactose and glucose for use in the food processing industry, the pharmaceutical industry, for example, to treat intolerance to lactose, as a diagnostic reporter molecule, in corn wet milling, in the fruit juice industry, in baking, in the textile industry and in the detergent industry.

In accordance with yet a further aspect of the present invention, there is provided a process for utilizing such enzymes for hydrolyzing guar gum (a galactomannan polysaccharide) to remove non-reducing terminal mannose residues. Further polysaccharides such as galactomannan and the enzymes according to the invention that degrade them have a variety of applications. Guar gum is commonly used as a thickening agent in food and also is utilized in hydraulic fracturing in oil and gas recovery. Consequently, mannanases are industrially relevant for the degradation and modification of guar gums. Furthermore, a need exists for thermostable mannases that are active in extreme conditions associated with drilling and well stimulation.

In accordance with yet a further aspect of the present invention, there are also provided nucleic acid probes comprising nucleic acid molecules of sufficient length to specifically hybridize to a nucleic acid sequence of the present invention.

In accordance with yet a further aspect of the present invention, there is provided process for utilizing such enzymes, or polynucleotides encoding such enzymes, for in vitro purposes related to scientific research, for example, to generate probes for identifying similar sequences which might encode similar enzymes from other organisms by using certain regions, i.e., conserved sequence regions, of the nucleotide sequence.

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These and other aspects of the present invention should be apparent to those skilled in the art from the teachings herein.

The polynucleotides of this invention were originally recovered from genomic gene libraries derived from the following organisms:

M11TL is a new species of *Desulfurococcus* isolated from Diamond Pool in Yellowstone National Park. The organism grows optimally at 85-88°C, pH 7.0 in a low salt medium containing yeast extract, peptone, and gelatin as substrates with a N₂/CO₂ gas phase.

OC1/4V is from the genus *Thermotoga*. The organism was isolated from Yellowstone National Park. It grows optimally at 75° C in a low salt medium with cellulose as a substrate and N_2 in gas phase.

Pyrococcus furiosus VC1 and (7EG1) is from the genus Pyrococcus. VC1 was isolated from Vulcano, Italy. It grows optimally at 100° C in a high salt medium (marine) containing elemental sulfur, yeast extract, peptone and starch as substrates and N_2 in gas phase.

Staphylothermus marinus F1 is a from the genus Staphylothermus. F1 was isolated from Vulcano, Italy. It grows optimally at 85°C, pH 6.5 in high salt medium (marine) containing elemental sulfur and yeast extract as substrates and N₂ in gas phase.

Thermococcus 9N-2 is from the genus Thermococcus 9N-2 was isolated from diffuse vent fluid in the East Pacific Rise. It is a strict anaerobe that grows optimally at 87°C.

Thermotoga maritima MSB8 and MSB8 (Clone # 6GP2 and 6GB4) is from the genus Thermotogo, and was isolated from Vulcano, Italy. MSB8 grows optimally at 85°C, pH 6.5 in a high salt medium (marine) containing starch and yeast extract as substrates and N₂ in gas phase.

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Thermococcus alcaliphilus AEDII12RA is from the genus Thermococcus. AEDII12RA grows optimally at 85°C, pH 9.5 in a high salt medium (marine) containing polysulfides and yeast extract as substrates and N_2 in gas phase.

Thermococcus chitonophagus GC74 is from the genus Thermococcus. GC74 grows optimally at 85°C, pH 6.0 in a high salt medium (marine) containing chitin, meat extract, elemental sulfur and yeast extract as substrates and N₂ in gas phase. AEPII 1a grows optimally at 85°C at pH 6.5 in marine medium under anaerobic conditions. It has many substrates. Bankia gouldi is from the genus Bankia.

Accordingly, the polynucleotides and enzymes encoded thereby are identified by the organism from which they were isolated, and are sometimes hereinafter referred to as "M11TL" (Figure 1 and SEQ ID NOS:1 and 15), "OC1/4V-33B/G" (Figure 2 and SEQ ID NOS:2 and 16), "F1-12G" (Figure 3 and SEQ ID NOS:3 and 17), "9N2-31B/G" (Figure 4 and SEQ ID NOS:4 and 18), "MSB8" (Figure 5 and SEQ ID NOS:5 and 19), "AEDII12RA-18B/G" (Figure 6 and SEQ ID NOS:6 and 20), "GC74-22G" (Figure 7 and SEQ ID NOS:7 and 21), "VC1-7G1" (Figure 8 and SEQ ID NOS:8 and 22), "37GP1" (Figure 9 and SEQ ID NOS: 9 and 23), "6GC2" (Figure 10 and SEQ ID NOS: 10 and 24), "6GP2" (Figure 11 and SEQ ID NOS:11 and 25), "AEPII 1a" (Figure 12 and SEQ ID NOS:12 and 26), "OC1/4V" (Figure 13 and SEQ ID NOS:13 and 27), and "6GP3" (Figure 14 and SEQ ID NOS:28), "MSB8-6GP2" (Figure 15 and SEQ ID NOS:57 and 61), "MSB8-6GB4"(Figure 16 and SEQ ID NOS:58 and 62),"VC1-7EG1"(Figure 17 and SEQ ID NOS:59 and 63), and 37GP4 (Figure 18 and SEQ ID NOS:60 and 64).

The polynucleotides and polypeptides of the present invention show identity at the nucleotide and protein level to known genes and proteins encoded thereby as shown in Table 1.

Table 1

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Clone	Gene/Protein with Closest Homology	Protein Identity	Nucleic Acid Identity
M11TL-29G	Sulfolobus sulfataricus DSM 1616/P1, β- galactosidase	51%	55%
OC1/4V-33B/G	Caldocellum saccharolyticum, β- glucosidase	52%	57%
Staphylothermus marinus F1-12G	Bacillus polymyxa, β- galactosidase	36%	48%
Thermococcus 9N2- 31B/G	Sulfolobus sulfataricus ATCC 49255/MT4, β- galactosidase	51%	50%
Thermotoga maritima MSB8-6G	Clostridium thermocellum	45%	53%
Thermococcus AEDII12RA-18B/G	Bacillus polymyxa, β- galactosidase	34%	48%
Thermococcus chitonophagus GC74- 22G	Sulfolobus sulfataricus ATCC 49255/MT4, β- galactosidase	46%	54%

Pyrococcus furiosus	Sulfolobus	46.4%	52.5%
VC1-7G1	sulfataricus/MT-4 β-		
•	galactosidase		
Thermotoga maritima	Pediococcus rentosaceaus	49%	29%
m-galactosidase	α-galactosidase		
(6GC2)			
Thermotoga maritima	Aspergillus aculeatus	56%	37%
ß-mannanase (6GP2)	mannanase		
AEPII 1a ß-	Sulfolobus solfactaricus ß-	78%	56%
mannosidase (63GB1)	galactosidase		· · · · · · · · · · · · · · · · · · ·
OC1/4V	Clostridium thermocellum	65%	43%
endoglucanase	endo-1,4-ß-endoglucanase		
(33GP1)			
Thermotoga maritima	Caldocellum	72	53
pullalanase (6GP3)	saccharolyticum α-		•
	destrom 6		
	glucanohydralase		
Bankia gouldi mix	None available		
Endoglucanase			
(37GP1)			

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The polynucleotides and enzymes of the present invention show homology to each other as shown in Table 2.

Table 2

Clone	Gene/Protein with Closest Homology	Protein Identity	Nucleic Acid Identity
Staphylothermus marinus F1-12G	Thermococcus AEDII12RA-18B/G, β- galactosidase, glucosidase	55%	57%
Thermococcus 9N2- 31B/G	Thermococcus chitonophagus GC74- 22G-glucosidase`	74%	66%
Pyrococcus furiosus VC1-7G1	Pyrococcus furiosus VC1- 7B/G β-galactosidase	46.4%	54%

All the clones identified in Tables 1 and 2 encode polypeptides which have α -glycosidase or β -glycosidase activity.

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This invention, in addition to the isolated nucleic acid molecules encoding the enzymes of the present invention, also provide substantially similar sequences. Isolated nucleic acid sequences are substantially similar if: (i) they are capable of hybridizing under conditions hereinafter described, to the polynucleotides of SEQ ID NOS: 1-14 and 57-60; (ii) or they encode DNA sequences which are degenerate to the polynucleotides of SEQ ID NOS: 1-14 and 57-60. Degenerate DNA sequences encode the amino acid sequences of SEQ ID NOS:15-28 and 61-64, but have variations in the nucleotide coding sequences. As used herein, substantially similar refers to the sequences having similar identity to the sequences of the instant invention. The nucleotide sequences that are substantially the same can be identified by hybridization or by sequence comparison. Enzyme sequences that are substantially the same can be identified by one or more of the following: proteolytic digestion, gel electrophoresis and/or microsequencing.

One means for isolating the nucleic acid molecules encoding the enzymes of the present invention is to probe a gene library with a natural or artificially designed probe using art recognized procedures (see, for example: Current Protocols in Molecular Biology,

Ausubel F.M. et al. (EDS.) Green Publishing Company Assoc. and John Wiley Interscience, New York, 1989, 1992). It is appreciated to one skilled in the art that the polynucleotides of SEQ ID NOS: 1-14 and 57-60 or fragments thereof (comprising at least 12 contiguous nucleotides), are particularly useful probes. Other particular useful probes for this purpose are hybridizable fragments to the sequences of SEQ ID NOS: 1-14 and 57-60 (i.e., comprising at least 12 contiguous nucleotides).

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With respect to nucleic acid sequences which hybridize to specific nucleic acid sequences disclosed herein, hybridization may be carried out under conditions of reduced stringency, medium stringency or even stringent conditions. As an example of oligonucleotide hybridization, a polymer membrane containing immobilized denatured nucleic acids is first prehybridized for 30 minutes at 45 °C in a solution consisting of 0.9 M NaCl, 50 mM NaH₂PO₄, pH 7.0, 5.0 mM Na₂EDTA, 0.5% SDS, 10X Denhardt's, and 0.5 mg/ml polyriboadenylic acid. Approximately 2 X 10⁷ cpm (specific activity 4-9 X 10 cpm/ug) of ³²P end-labeled oligonucleotide probe are then added to the solution. After 12-16 hours of incubation, the membrane is washed for 30 minutes at room temperature in 1X SET (150 mM NaCl, 20 mM Tris hydrochloride, pH 7.8, 1 mM Na₂EDTA) containing 0.5% SDS, followed by a 30 minute wash in fresh 1X SET at Tm 10 °C for the oligonucleotide probe. The membrane is then exposed to auto-radiographic film for detection of hybridization signals.

Stringent conditions means hybridization will occur only if there is at least 90% identity, preferably at least 95% identity and most preferably at least 97% identity between the sequences. Further, it is understood that a section of a 100 bps sequence that is 95 bps in length has 95% identity with the 1090 bps sequence from which it is obtained. See J. Sambrook et al., Molecular Cloning, A Laboratory Manual, 2d Ed., Cold Spring Harbor Laboratory (1989) which is hereby incorporated by reference in its entirety. Also, it is understood that a fragment of a 100 bps sequence that is 95 bps in length has 95% identity with the 100 bps sequence from which it is obtained.

As used herein, a first DNA (RNA) sequence is at least 70% and preferably at least 80% identical to another DNA (RNA) sequence if there is at least 70% and preferably at

least a 80% or 90% identity, respectively, between the bases of the first sequence and the bases of the another sequence, when properly aligned with each other, for example when aligned by BLASTN.

"Identity" as the term is used herein, refers to a polynucleotide sequence which comprises a percentage of the same bases as a reference polynucleotide (SEQ ID NOS:1-14 and 57-60). For example, a polynucleotide which is at least 90% identical to a reference polynucleotide, has polynucleotide bases which are identical in 90% of the bases which make up the reference polynucleotide and may have different bases in 10% of the bases which comprise that polynucleotide sequence.

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The present invention relates polynucleotides which differ from the reference polynucleotide such that the changes are silent changes, for example the change do not alter the amino acid sequence encoded by the polynucleotide. The present invention also relates to nucleotide changes which result in amino acid substitutions, additions, deletions, fusions and truncations in the polypeptide encoded by the reference polynucleotide. In a preferred aspect of the invention these polypeptides retain the same biological action as the polypeptide encoded by the reference polynucleotide.

It is also appreciated that such probes can be and are preferably labeled with an analytically detectable reagent to facilitate identification of the probe. Useful reagents include but are not limited to radioactivity, fluorescent dyes or enzymes capable of catalyzing the formation of a detectable product. The probes are thus useful to isolate complementary copies of DNA from other sources or to screen such sources for related sequences.

The polynucleotides of this invention were recovered from genomic gene libraries from the organisms listed in Table 1. For example, gene libraries can be generated in the Lambda ZAP II cloning vector (Stratagene Cloning Systems). Mass excisions can be performed on these libraries to generate libraries in the pBluescript phagemid. Libraries are thus generated and excisions performed according to the protocols/methods hereinafter described.

The excision libraries are introduced into the *E. coli* strain BW14893 F'kan1A. Expression clones are then identified using a high temperature filter assay. Expression clones encoding several glucanases and several other glycosidases are identified and repurified. The polynucleotides, and enzymes encoded thereby, of the present invention, yield the activities as described above.

The coding sequences for the enzymes of the present invention were identified by screening the genomic DNAs prepared for the clones having glucosidase or galactosidase activity.

An example of such an assay is a high temperature filter assay wherein expression clones were identified by use of high temperature filter assays using buffer Z (see recipe below) containing 1 mg/ml of the substrate 5-bromo-4-chloro-3-indolyl-β-D-glucopyranoside (XGLU) (Diagnostic Chemicals Limited or Sigma) after introducing an excision library into the *E. coli* strain BW14893 F'kan1A. Expression clones encoding XGLUases were identified and repurified from M11TL, OC1/4V, Pyrococcus furiosus VC1, Staphylothemus marinus F1. Thermococcus 9N-2, Thermotoga maritima MSB8, Thermococcus alcaliphilus AEDII12RA, and Thermococcus chitonophagus GC74.

Z-buffer: (referenced in Miller, J.H. (1992) A Short Course in Bacterial Genetics, p. 445.)

per liter:

Adjust pH to 7.0

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 $Na_2HPO_4-7H_2O$ 16.1g $NaH_2PO_4-7H_2O$ 5.5g KCl 0.75g $MgSO_4-7H_2O$ 0.246g β-mercaptoethanol 2.7ml

High Temperature Filter Assay

(1) The f factor fkan (from E. coli strain CSH118)(1) was introduced into the pho-pnh-lac-strain BW14893(2). BW13893(2). The filamentous phage library was plated on the resulting strain, BW14893 F'kan. (Miller, J.H. (1992) A Short Course in

Bacterial Genetics, Lee, K.S., Metcalf, et al., (1992) Evidence for two phosphonate degradative pathways in Enterobacter Aerogenes, J. Bacteriol., 174:2501-2510.

(2) After growth on 100 mm LB plates containing 100 μg/ml ampicillin, 80 μg/ml nethicillin and 1mM IPTG, colony lifts were performed using Millipore HATF membrane filters.

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- (3) The colonies transferred to the filters were lysed with chloroform vapor in 150 mm glass petri dishes.
- (4) The filters were transferred to 100 mm glass petri dishes containing a piece of Whatman 3MM filter paper saturated with buffer.
 - (a) when testing for galactosidase activity (XGALase), 3MM paper was saturated with Z buffer containing 1 mg/ml XGAL (ChemBridge Corporation). After transferring filter bearing lysed colonies to the glass petri dish, placed dish in oven at 80-85°C.
 - (b) when testing for glucosidase (XGLUase), 3MM paper was saturated with Z buffer containing 1 mg/ml XGLU. After transferring filter bearing lysed colonies to the glass petri dish, placed dish in oven at 80-85°C.
- (5) 'Positives' were observed as blue spots on the filter membranes. Used the following filter rescue technique to retrieve plasmid from lysed positive colony. Used pasteur pipette (or glass capillary tube) to core blue spots on the filter membrane. Placed the small filter disk in an Eppendorf tube containing 20 μl water. Incubated the Eppendorf tube at 75°C for 5 minutes followed by vortexing to elute plasmid DNA off filter. This DNA was transformed into electrocompetent *E. coli* cells DH10B for Thermatoga maritima MSB8-6G, Staphylothermus marinus F1-12G, Thermococcus AEDII12RA-18B/G, Thermococcus chitonophagus GC74-22G, M11Tl and OC1/4V. Electrocompetent BW14893 F'kan1A *E. coli* were used for Thermococcus 9N2-31B/G, and *Pyrococcus furiosus* VC1-7G1. Repeated filter-lift assay on transformation plates to identify 'positives'. Return transformation plates to 37°C incubator after filter lift to regenerate colonies. Inoculate 3 ml LB liquid containing 100 μg/ml ampicillin with repurified positives and incubate at 37°C

overnight. Isolate plasmid DNA from these cultures and sequence plasmid insert. In some instances where the plates used for the initial colony lifts contained non-confluent colonies, a specific colony corresponding to a blue spot on the filter could be identified on a regenerated plate and repurified directly, instead of using the filter rescue technique.

Another example of such an assay is a variation of the high temperature filter assay wherein colony-laden filters are heat-killed at different temperatures (for example, 105°C for 20 minutes) to monitor thermostability. The 3MM paper is saturated with different buffers (i.e., 100 mM NaCl, 5 mM MgCl₂, 100 mM Tris-Cl (pH 9.5)) to determine enzyme activity under different buffer conditions.

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A β -glucosidase assay may also be employed, wherein Glcp β Np is used as an artificial substrate (aryl- β -glucosidase). The increase in absorbance at 405 nm as a result of p-nitrophenol (pNp) liberation was followed on a Hitachi U-1100 spectrophotometer, equipped with a thermostatted cuvette holder. The assays may be performed at 80°C or 90°C in closed 1-ml quartz cuvette. A standard reaction mixture contains 150 mM trisodium substrate, pH 5.0 (at 80°C), and 0.95 mM pNp derivative pNp = 0.561 mM⁻¹ cm⁻¹). The reaction mixture is allowed to reach the desired temperature, after which the reaction is started by injecting an appropriate amount of enzyme (1.06 ml final volume).

1 U β -glucosidase activity is defined as that amount required to catalyze the formation of 1.0 μ mol pNp/min. D-cellobiose may also be used as a substrate.

An ONPG assay for β -galactosidase activity is described by Miller, J.H. (1992) A Short Course in Bacterial Genetics and Mill, J.H. (1992) Experiments in Molecular Genetics, the contents of which are hereby incorporated by reference in their entirety.

A quantitative fluorometric assay for β -galactosidase specific activity is described by : Youngman P., (1987) Plasmid Vectors for Recovering and Exploiting Tn917 Transpositions in Bacillus and other Gram-Positive Bacteria. In Plasmids: A Practical approach (ed. K. Hardy) pp 79-103. IRL Press, Oxford. A description of the procedure can be found in Miller (1992) p. 75-77, the contents of which are incorporated by reference herein in their entirety.

The polynucleotides of the present invention may be in the form of DNA which DNA includes cDNA, genomic DNA, and synthetic DNA. The DNA may be double-stranded or single-stranded, and if single stranded may be the coding strand or non-coding (anti-sense) strand. The coding sequences which encodes the mature enzymes may be identical to the coding sequences shown in Figures 1-8 (SEQ ID NOS: 1-14 and 57-60) or may be a different coding sequence which coding sequence, as a result of the redundancy or degeneracy of the genetic code, encodes the same mature enzymes as the DNA of Figures 1-18 (SEQ ID NOS: 1-14 and 57-60).

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The polynucleotide which encodes for the mature enzyme of Figures 1-18 (SEQ ID NOS: 15-28 and 61-64) may include, but is not limited to: only the coding sequence for the mature enzyme; the coding sequence for the mature enzyme and additional coding sequence such as a leader sequence or a proprotein sequence; the coding sequence for the mature enzyme (and optionally additional coding sequence) and non-coding sequence, such as introns or non-coding sequence 5' and/or 3' of the coding sequence for the mature enzyme.

Thus, the term "polynucleotide encoding an enzyme (protein)" encompasses a polynucleotide which includes only coding sequence for the enzyme as well as a polynucleotide which includes additional coding and/or non-coding sequence.

The present invention further relates to variants of the hereinabove described polynucleotides which encode for fragments, analogs and derivatives of the enzymes having the deduced amino acid sequences of Figures 1-18 (SEQ ID NOS: 15-28 and 61-64). The variant of the polynucleotide may be a naturally occurring allelic variant of the polynucleotide or a non-naturally occurring variant of the polynucleotide.

Thus, the present invention includes polynucleotides encoding the same mature enzymes as shown in Figures 1-18 (SEQ ID NOS: 15-28 and 61-64) as well as variants of such polynucleotides which variants encode for a fragment, derivative or analog of the enzymes of Figures 1-18 (SEQ ID NOS: 15-28 and 61-64). Such nucleotide variants include deletion variants, substitution variants and addition or insertion variants.

As hereinabove indicated, the polynucleotides may have a coding sequence which is a naturally occurring allelic variant of the coding sequences shown in Figures 1-18 (SEQ

ID NOS: 1-14 and 57-60). As known in the art, an allelic variant is an alternate form of a polynucleotide sequence which may have a substitution, deletion or addition of one or more nucleotides, which does not substantially alter the function of the encoded enzyme.

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Fragments of the full length gene of the present invention may be used as a hybridization probe for a cDNA or a genomic library to isolate the full length DNA and to isolate other DNAs which have a high sequence similarity to the gene or similar biological activity. Probes of this type preferably have at least 10, preferably at least 15, and even more preferably at least 30 bases and may contain, for example, at least 50 or more bases. The probe may also be used to identify a DNA clone corresponding to a full length transcript and a genomic clone or clones that contain the complete gene including regulatory and promotor regions, exons, and introns. An example of a screen comprises isolating the coding region of the gene by using the known DNA sequence to synthesize an oligonucleotide probe. Labeled oligonucleotides having a sequence complementary to that of the gene of the present invention are used to screen a library of genomic DNA to determine which members of the library the probe hybridizes to.

The present invention further relates to polynucleotides which hybridize to the hereinabove-described sequences if there is at least 70%, preferably at least 90%, and more preferably at least 95% identity between the sequences. The present invention particularly relates to polynucleotides which hybridize under stringent conditions to the hereinabove-described polynucleotides. As herein used, the term "stringent conditions" means hybridization will occur only if there is at least 95% and preferably at least 97% identity between the sequences. The polynucleotides which hybridize to the hereinabove described polynucleotides in a preferred embodiment encode enzymes which either retain substantially the same biological function or activity as the mature enzyme encoded by the DNA of Figures 1-18 (SEQ ID NOS: 1-14 and 57-60).

Alternatively, the polynucleotide may have at least 15 bases, preferably at least 30 bases, and more preferably at least 50 bases which hybridize to any part of a polynucleotide of the present invention and which has an identity thereto, as hereinabove described, and which may or may not retain activity. For example, such polynucleotides may be employed

as probes for the polynucleotides of SEQ ID NOS: 1-14 and 57-60, for example, for recovery of the polynucleotide or as a diagnostic probe or as a PCR primer.

Thus, the present invention is directed to polynucleotides having at least a 70% identity, preferably at least 90% identity and more preferably at least a 95% identity to a polynucleotide which encodes the enzymes of SEQ ID NOS: 15-28 and 61-64 as well as fragments thereof, which fragments have at least 15 bases, preferably at least 30 bases and most preferably at least 50 bases, which fragments are at least 90% identical, preferably at least 95% identical and most preferably at least 97% identical under stringent conditions to any portion of a polynucleotide of the present invention.

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The present invention further relates to enzymes which have the deduced amino acid sequences of Figures 1-18 (SEQ ID NOS: 15-28 and 61-64) as well as fragments, analogs and derivatives of such enzyme.

The terms "fragment," "derivative" and "analog" when referring to the enzymes of Figures 1-18 (SEQ ID NOS: 15-28 and 61-64) means enzymes which retain essentially the same biological function or activity as such enzymes. Thus, an analog includes a proprotein which can be activated by cleavage of the proprotein portion to produce an active mature enzyme.

The enzymes of the present invention may be a recombinant enzyme, a natural enzyme or a synthetic enzyme, preferably a recombinant enzyme.

The fragment, derivative or analog of the enzymes of Figures 1-18 (SEQ ID NOS: 15-28 and 61-64) may be (i) one in which one or more of the amino acid residues are substituted with a conserved or non-conserved amino acid residue (preferably a conserved amino acid residue) and such substituted amino acid residue may or may not be one encoded by the genetic code, or (ii) one in which one or more of the amino acid residues includes a substituent group, or (iii) one in which the mature enzyme is fused with another compound, such as a compound to increase the half-life of the enzyme (for example, polyethylene glycol), or (iv) one in which the additional amino acids are fused to the mature enzyme, such as a leader or secretory sequence or a sequence which is employed for purification of the mature enzyme or a proprotein sequence. Such fragments, derivatives

and analogs are deemed to be within the scope of those skilled in the art from the teachings herein.

The enzymes and polynucleotides of the present invention are preferably provided in an isolated form, and preferably are purified to homogeneity.

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The term "isolated" means that the material is removed from its original environment (e.g., the natural environment if it is naturally occurring). For example, a naturally-occurring polynucleotide or enzyme present in a living animal is not isolated, but the same polynucleotide or enzyme, separated from some or all of the coexisting materials in the natural system, is isolated. Such polynucleotides could be part of a vector and/or such polynucleotides or enzymes could be part of a composition, and still be isolated in that such vector or composition is not part of its natural environment.

The enzymes of the present invention include the enzymes of SEQ ID NOS: 15-28 and 61-64 (in particular the mature enzyme) as well as enzymes which have at least 70% similarity (preferably at least 70% identity) to the enzymes of SEQ ID NOS: 15-28 and 61-64 and more preferably at least 90% similarity (more preferably at least 90% identity) to the enzymes of SEQ ID NOS: 15-28 and 61-64 and still more preferably at least 95% similarity (still more preferably at least 95% identity) to the enzymes of SEQ ID NOS: 15-28 and 61-64 and also include portions of such enzymes with such portion of the enzyme generally containing at least 30 amino acids and more preferably at least 50 amino acids.

As known in the art "similarity" between two enzymes is determined by comparing the amino acid sequence and its conserved amino acid substitutes of one enzyme to the sequence of a second enzyme.

A variant, i.e. a "fragment", "analog" or "derivative" polypeptide, and reference polypeptide may differ in amino acid sequence by one or more substitutions, additions, deletions, fusions and truncations, which may be present in any combination.

Among preferred variants are those that vary from a reference by conservative amino acid substitutions. Such substitutions are those that substitute a given amino acid in a polypeptide by another amino acid of like characteristics. Typically seen as conservative substitutions are the replacements, one for another, among the aliphatic amino acids Ala,

Val, Leu and Ile; interchange of the hydroxyl residues Ser and Thr, exchange of the acidic residues Asp and Glu, substitution between the amide residues Asn and Gln, exchange of the basic residues Lys and Arg and replacements among the aromatic residues Phe, Tyr.

Most highly preferred are variants which retain the same biological function and activity as the reference polypeptide from which it varies.

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Fragments or portions of the enzymes of the present invention may be employed for producing the corresponding full-length enzyme by peptide synthesis; therefore, the fragments may be employed as intermediates for producing the full-length enzymes. Fragments or portions of the polynucleotides of the present invention may be used to synthesize full-length polynucleotides of the present invention.

The present invention also relates to vectors which include polynucleotides of the present invention, host cells which are genetically engineered with vectors of the invention and the production of enzymes of the invention by recombinant techniques.

Host cells are genetically engineered (transduced or transformed or transfected) with the vectors of this invention which may be, for example, a cloning vector or an expression vector. The vector may be, for example, in the form of a plasmid, a viral particle, a phage, etc. The engineered host cells can be cultured in conventional nutrient media modified as appropriate for activating promoters, selecting transformants or amplifying the genes of the present invention. The culture conditions, such as temperature, pH and the like, are those previously used with the host cell selected for expression, and will be apparent to the ordinarily skilled artisan.

The polynucleotides of the present invention may be employed for producing enzymes by recombinant techniques. Thus, for example, the polynucleotide may be included in any one of a variety of expression vectors for expressing an enzyme. Such vectors include chromosomal, nonchromosomal and synthetic DNA sequences, e.g., derivatives of SV40; bacterial plasmids; phage DNA; baculovirus; yeast plasmids; vectors derived from combinations of plasmids and phage DNA, viral DNA such as vaccinia, adenovirus, fowl pox virus, and pseudorabies. However, any other vector may be used as long as it is replicable and viable in the host.

The appropriate DNA sequence may be inserted into the vector by a variety of procedures. In general, the DNA sequence is inserted into an appropriate restriction endonuclease site(s) by procedures known in the art. Such procedures and others are deemed to be within the scope of those skilled in the art.

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The DNA sequence in the expression vector is operatively linked to an appropriate expression control sequence(s) (promoter) to direct mRNA synthesis. As representative examples of such promoters, there may be mentioned: LTR or SV40 promoter, the <u>E. coli</u>. lac or trp, the phage lambda P_L promoter and other promoters known to control expression of genes in prokaryotic or eukaryotic cells or their viruses. The expression vector also contains a ribosome binding site for translation initiation and a transcription terminator. The vector may also include appropriate sequences for amplifying expression.

In addition, the expression vectors preferably contain one or more selectable marker genes to provide a phenotypic trait for selection of transformed host cells such as dihydrofolate reductase or neomycin resistance for eukaryotic cell culture, or such as tetracycline or ampicillin resistance in <u>E. coli</u>.

The vector containing the appropriate DNA sequence as hereinabove described, as well as an appropriate promoter or control sequence, may be employed to transform an appropriate host to permit the host to express the protein.

As representative examples of appropriate hosts, there may be mentioned: bacterial cells, such as <u>E. coli</u>, <u>Streptomvces</u>, <u>Bacillus subtilis</u>; fungal cells, such as yeast; insect cells such as <u>Drosophila S2</u> and <u>Spodoptera Sf9</u>; animal cells such as CHO, COS or Bowes melanoma; adenoviruses; plant cells, etc. The selection of an appropriate host is deemed to be within the scope of those skilled in the art from the teachings herein.

More particularly, the present invention also includes recombinant constructs comprising one or more of the sequences as broadly described above. The constructs comprise a vector, such as a plasmid or viral vector, into which a sequence of the invention has been inserted, in a forward or reverse orientation. In a preferred aspect of this embodiment, the construct further comprises regulatory sequences, including, for example, a promoter, operably linked to the sequence. Large numbers of suitable vectors and

promoters are known to those of skill in the art. and are commercially available. The following vectors are provided by way of example; Bacterial: pQE70, pQE60, pQE-9 (Qiagen), pD10, psiX174, pBluescript iI KS, pNH8A, pNH16a, pNH18A, pNH46A (Stratagene); ptrc99a, pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia); Eukaryotic: pSV2CAT, pOG44, pXT1, pSG (Stratagene) pSVK3, pBPV, pMSG, pSVL (Pharmacia). However, any other plasmid or vector may be used as long as they are replicable and viable in the host.

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Promoter regions can be selected from any desired gene using CAT (chloramphenical transferase) vectors or other vectors with selectable markers. Two appropriate vectors are pKK232-8 and pCM7. Particular named bacterial promoters include lacI, lacZ, T3, T7, gpt, lambda P_R, P_L and trp. Eukaryotic promoters include CMV immediate early, HSV thymidine kinase, early and late SV40, LTRs from retrovirus, and mouse metallothionein-I. Selection of the appropriate vector and promoter is well within the level of ordinary skill in the art.

In a further embodiment, the present invention relates to host cells containing the above-described constructs. The host cell can be a higher eukaryotic cell, such as a mammalian cell, or a lower eukaryotic cell, such as a yeast cell, or the host cell can be a prokaryotic cell, such as a bacterial cell. Introduction of the construct into the host cell can be effected by calcium phosphate transfection, DEAE-Dextran mediated transfection, or electroporation (Davis, L., Dibner, M., Battey, I., Basic Methods in Molecular Biology, (1986)).

The constructs in host cells can be used in a conventional manner to produce the gene product encoded by the recombinant sequence. Alternatively, the enzymes of the invention can be synthetically produced by conventional peptide synthesizers.

Mature proteins can be expressed in mammalian cells, yeast, bacteria, or other cells under the control of appropriate promoters. Cell-free translation systems can also be employed to produce such proteins using RNAs derived from the DNA constructs of the present invention. Appropriate cloning and expression vectors for use with prokaryotic and eukaryotic hosts are described by Sambrook, et al., Molecular Cloning: A Laboratory

Manual, Second Edition, Cold Spring Harbor, N.Y., (1989), the disclosure of which is hereby incorporated by reference.

Transcription of the DNA encoding the enzymes of the present invention by higher eukaryotes is increased by inserting an enhancer sequence into the vector. Enhance, are cis-acting elements of DNA, usually about from 10 to 300 bp that act on a promoter to increase its transcription. Examples include the SV40 enhancer on the late side of the replication origin bp 100 to 270, a cytomegalovirus early promoter enhancer, the polyoma enhancer on the late side of the replication origin, and adenovirus enhancers.

Generally, recombinant expression vectors will include origins of replication and selectable markers permitting transformation of the host cell, e.g., the ampicillin resistance gene of E. coli and S. cerevisiae TRP1 gene, and a promoter derived from a highly-expressed gene to direct transcription of a downstream structural sequence. Such promoters can be derived from operons encoding glycolytic enzymes such as 3-phosphoglycerate kinase (PGK), α-factor, acid phosphatase, or heat shock proteins, among others. The heterologous structural sequence is assembled in appropriate phase with translation initiation and termination sequences, and preferably, a leader sequence capable of directing secretion of translated enzyme. Optionally, the heterologous sequence can encode a fusion enzyme including an N-terminal identification peptide imparting desired characteristics, e.g., stabilization or simplified purification of expressed recombinant product.

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Useful expression vectors for bacterial use are constructed by inserting a structural DNA sequence encoding a desired protein together with suitable translation initiation and termination signals in operable reading phase with a functional promoter. The vector will comprise one or more phenotypic selectable markers and an origin of replication to ensure maintenance of the vector and to, if desirable, provide amplification within the host. Suitable prokaryotic hosts for transformation include <u>E. coli, Bacillus subtilis, Salmonella typhimurium</u> and various species within the genera Pseudomonas, Streptomyces, and Staphylococcus, although others may also be employed as a matter of choice.

As a representative but nonlimiting example, useful expression vectors for bacterial use can comprise a selectable marker and bacterial origin of replication derived from

commercially available plasmids comprising genetic elements of the well known cloning vector pBR322 (ATCC 37017). Such commercial vectors include, for example, pKK223-3 (Pharmacia Fine Chemicals, Uppsala, Sweden) and GEM1 (Promega Biotec, Madison, WI, USA). These pBR322 "backbone" sections are combined with an appropriate promoter and the structural sequence to be expressed.

Following transformation of a suitable host strain and growth of the host strain to an appropriate cell density, the selected promoter is induced by appropriate means (e.g., temperature shift or chemical induction) and cells are cultured for an additional period.

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Cells are typically harvested by centrifugation, disrupted by physical or chemical means, and the resulting crude extract retained for further purification.

Microbial cells employed in expression of proteins can be disrupted by any convenient method, including freeze-thaw cycling, sonication, mechanical disruption, or use of cell lysing agents, such methods are well known to those skilled in the art.

Various mammalian cell culture systems can also be employed to express recombinant protein. Examples of mammalian expression systems include the COS-7 lines of monkey kidney fibroblasts, described by Gluzman, Cell, 23:175 (1981), and other cell lines capable of expressing a compatible vector, for example, the C127, 3T3, CHO, HeLa and BHK cell lines. Mammalian expression vectors will comprise an origin of replication, a suitable promoter and enhancer, and also any necessary ribosome binding sites, polyadenylation site, splice donor and acceptor sites, transcriptional termination sequences, and 5' flanking nontranscribed sequences. DNA sequences derived from the SV40 splice, and polyadenylation sites may be used to provide the required nontranscribed genetic elements.

The enzyme can be recovered and purified from recombinant cell cultures by methods including ammonium sulfate or ethanol precipitation, acid extraction, anion or cation exchange chromatography, phosphocellulose chromatography, hydrophobic interaction chromatography, affinity chromatography, hydroxylapatite chromatography and lectin chromatography. Protein refolding steps can be used, as necessary, in completing

configuration of the mature protein. Finally, high performance liquid chromatography (HPLC) can be employed for final purification steps.

The enzymes of the present invention may be a naturally purified product, or a product of chemical synthetic procedures, or produced by recombinant techniques from a prokaryotic or eukaryotic host (for example, by bacterial, yeast, higher plant, insect and mammalian cells in culture). Depending upon the host employed in a recombinant production procedure, the enzymes of the present invention may be glycosylated or may be non-glycosylated. Enzymes of the invention may or may not also include an initial methionine amino acid residue.

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 β -galactosidase hydrolyzes lactose to galactose and glucose. Accordingly, the OC1/4V, 9N2-31B/G, AEDII12RA-18B/G and F1-12G enzymes may be employed in the food processing industry for the production of low lactose content milk and for the production of galactose or glucose from lactose contained in whey obtained in a large amount as a by-product in the production of cheese. Generally, it is desired that enzymes used in food processing, such as the aforementioned β -galactosidases, be stable at elevated temperatures to help prevent microbial contamination.

These enzymes may also be employed in the pharmaceutical industry. The enzymes are used to treat intolerance to lactose. In this case, a thermostable enzyme is desired, as well. Thermostable β -galactosidases also have uses in diagnostic applications, where they are employed as reporter molecules.

Glucosidases act on soluble cellooligosaccharides from the non-reducing end to give glucose as the sole product. Glucanases (endo- and exo-) act in the depolymerization of cellulose, generating more non-reducing ends (endo-glucanases, for instance, act on internal linkages yielding cellobiose, glucose and cellooligosaccharides as products). β -glucosidases are used in applications where glucose is the desired product. Accordingly, M11TL, F1-12G, GC74-22G, MSB8-6G, OC1/4V, VC1-7G1, 9N2-31B/G and AEDII12RA18B/G may be employed in a wide variety of industrial applications, including in corn wet milling for the separation of starch and gluten, in the fruit industry for clarification and equipment maintenance, in baking for viscosity reduction, in the textile

industry for the processing of blue jeans, and in the detergent industry as an additive. For these and other applications, thermostable enzymes are desirable.

Antibodies generated against the etzymes corresponding to a sequence of the present invention can be obtained by direct injection of the enzymes into an animal or by administering the enzymes to an animal, preferably a nonhuman. The antibody so obtained will then bind the enzymes itself. In this manner, even a sequence encoding only a fragment of the enzymes can be used to generate antibodies binding the whole native enzymes. Such antibodies can then be used to isolate the enzyme from cells expressing that enzyme.

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For preparation of monoclonal antibodies, any technique which provides antibodies produced by continuous cell line cultures can be used. Examples include the hybridoma technique (Kohler and Milstein, 1975, Nature, 256:495-497), the trioma technique, the human B-cell hybridoma technique (Kozbor et al., 1983, Immunology Today 4:72), and the EBV-hybridoma technique to produce human monoclonal antibodies (Cole, et al., 1985, in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp. 77-96).

Techniques described for the production of single chain antibodies (U.S. Patent 4,946,778) can be adapted to produce single chain antibodies to immunogenic enzyme products of this invention. Also, transgenic mice may be used to express humanized antibodies to immunogenic enzyme products of this invention.

Antibodies generated against the enzyme of the present invention may be used in screening for similar enzymes from other organisms and samples. Such screening techniques are known in the art, for example, one such screening assay is described in "Methods for Measuring Cellulase Activities", *Methods in enzymology*, Vol 160, pp. 87-116, which is hereby incorporated by reference in its entirety.

The present invention will be further described with reference to the following examples; however, it is to be understood that the present invention is not limited to such examples. All parts or amounts, unless otherwise specified, are by weight.

In order to facilitate understanding of the following examples certain frequently occurring methods and/or terms will be described.

"Plasmids" are designated by a lower case p preceded and/or followed by capital letters and/or numbers. The starting plasmids herein are either commercially available, publicly available on an unrestricted basis, or can be constructed from available plasmids in accord with published procedures. In addition, equivalent plasmids to those described are known in the art and will be apparent to the ordinarily skilled artisan.

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"Digestion" of DNA refers to catalytic cleavage of the DNA with a restriction enzyme that acts only at certain sequences in the DNA. The various restriction enzymes used herein are commercially available and their reaction conditions, cofactors and other requirements were used as would be known to the ordinarily skilled artisan. For analytical purposes, typically 1 µg of plasmid or DNA fragment is used with about 2 units of enzyme in about 20 µl of buffer solution. For the purpose of isolating DNA fragments for plasmid construction, typically 5 to 50 µg of DNA are digested with 20 to 250 units of enzyme in a larger volume. Appropriate buffers and substrate amounts for particular restriction enzymes are specified by the manufacturer. Incubation times of about 1 hour at 37°C are ordinarily used, but may vary in accordance with the supplier's instructions. After digestion the reaction is electrophoresed directly on a polyacrylamide gel to isolate the desired fragment.

Size separation of the cleaved fragments is performed using 8 percent polyacrylamide gel described by Goeddel, D. et al., Nucleic Acids Res., 8:4057 (1980).

"Oligonucleotides" refers to either a single stranded polydeoxynucleotide or two complementary polydeoxynucleotide strands which may be chemically synthesized. Such synthetic oligonucleotides have no 5' phosphate and thus will not ligate to another oligonucleotide without adding a phosphate with an ATP in the presence of a kinase. A synthetic oligonucleotide will ligate to a fragment that has not been dephosphorylated.

"Ligation" refers to the process of forming phosphodiester bonds between two double stranded nucleic acid fragments (Maniatis, T., et al., Id., p. 146). Unless otherwise provided, ligation may be accomplished using known buffers and conditions with 10 units of T4 DNA ligase ("ligase") per 0.5 µg of approximately equimolar amounts of the DNA fragments to be ligated.

Unless otherwise stated, transformation was performed as described in the method of Graham, F. and Van der Eb, A., Virology, 52:456-457 (1973).

Example 1

Bacterial Expression and Purincation of Glycosidase Enzymes

DNA encoding the enzymes of the present invention, SEQ ID NOS: 1-14 and 57-60 were initially amplified from a pBluescript vector containing the DNA by the PCR technique using the primers noted herein. The amplified sequences were then inserted into the respective PQE vector listed beneath the primer sequences, and the enzyme was expressed according to the protocols set forth herein. The 5' and 3' primer sequences for the respective genes are as follows:

Thermococcus AEDII12RA -18B/G

5' CCGAGAATTCATTAAAGAGGAGAAATTAACTATGGTGAATGCTATGATTGTC 3' (SEQ ID NO:29)

3' CGGAAGATCTTCATAGCTCCGGAAGCCCATA 5' (SEQ ID NO:30)

Vector: pQE12; and contains the following restriction enzyme sites 5' EcoRI and 3' Blg II.

OC1/4V-33B/G

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5' CCGAGAATTCATTAAAGAGGAGAAATTAACTATGATAAGAAGGTCCGATTTTCC 3' (SEO ID NO:31)

3' CGGAAGATCTTTAAGATTTTAGAAATTCCTT 5' (SEQ ID NO:32)

Vector: pQE12; and contains the following restriction enzyme sites 5' EcoRI and 3' Bgl II.

Thermococcus 9N2 - 31B/G

5' CCGAGAATTCATTAAAGAGGGAGAAATTAACTATGCTACCAGAAGGCTTTCTC 3' (SEQ ID NO:33)

3' CGGAGGTACCTCACCCAAGTCCGAACTTCTC 5' (SEQ ID NO:34)

Vector: pQE30; and contains the following restriction enzyme sites 5' EcoRI and 3' KpnI.

Staphylothermus marinus F1 - 12G

5' CCGAGAATTCATTAAAGAGGAGAAATTAACTATGATAAGGTTTCCTGATTAT 3' (SEQ ID NO:35)

3' CGGAAGATCTTTATTCGAGGTTCTTTAATCC 5' (SEQ ID NO:36)

Vector: pQE12; and contains the following restriction enzyme sites 5' EcoRI and 3' Bgl II.

Thermococcus chitonophagus GC74 - 22G

5' CCGAGAATTCATTAAAGAGGAGAAATTAACTATGCTTCCAGGAGAACTTTCTC 3' (SEQ ID NO:37)

3' CGGAGGATCCCTACCCTCTCTAAGATCTC 5' (SEQ ID NO:38)

Vector: pQE12; and contains the following restriction enzyme sites 5' EcoRI and 3' BamHI.

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5' AATAATCTAGAGCATGCAATTCCCCAAAGACTTCATGATAG 3' (SEQ ID NO:39)

3' AATAAAAGCTTACTGGATCAGTGTAAGATGCT 5' (SEQ ID NO:40)

Vector: pQE70; and contains the following restriction enzyme sites 5' SphI and 3' Hind III.

Thermotoga maritima MSB8-6G

5' CCGACAATTGATTAAAGAGGAGAAATTAACTATGGAAAGGATCGATGAAATT 3' (SEQ ID NO:41)

3' CGGAGGTACCTCATGGTTTGAATCTCTTCTC 5' (SEQ ID NO:42)

Vector: pQE12; and contains the following restriction enzyme sites 5' EcoRI and 3' KpnI.

Pyrococcus furiosus VC1 - 7G1

5' CCGACAATTGATTAAAGAGGAGAAATTAACTATGTTCCCTGAAAAGTTCCTT 3' (SEQ ID NO:43)

3' CGGAGGTACCTCATCCCCTCAGCAATTCCTC 5' (SEQ ID NO:44)

Vector: pQE12; and contains the following restriction enzyme sites 5' EcoRI and 3' Kpn I.

Bankia gouldi endoglucanase (37GP1)

5' AATAAGGATCCGTTTAGCGACGCTCGC 3' (SEQ ID NO:45)

3' AATAAAAGCTTCCGGGTTGTACAGCGGTAATAGGC 5' (SEQ ID NO:46)

Vector: pQE52; and contains the following restriction enzyme sites 5' Bam HI and 3' Hind III.

Thermotoga maritima α-galactosidase (6GC2)

5' TTTATTGAATTCATTAAAGAGGAGAAATTAACTATGATCTGTGTGGAAATATTCGGAAAG 3' (SEQ ID NO:47)

3' TCTATAAAGCTTTCATTCTCTCACCCTCTTCGTAGAAG 5' (SEQ ID NO:48)

Vector: pQET; and contains the following restriction enzyme sites 5' EcoRI and 3' Hind III.

Thermotoga maritima \(\beta\)-mannanase (6GP2)

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5' TTTATTCAATTGATTAAAGAGGAGAAATTAACTATGGGGATTGGTGGCGACGAC 3' (SEQ ID NO:49)

3' TTTATTAAGCTTATCTTTTCATATTCACATACCTCC 5' (SEQ ID NO:50)

Vector: pQEt; and contains the following restriction enzyme sites 5' Hind III and 3' EcoRI.

AEPII Ia β-mannanase (63GB1)

5' TITATTGAATTCATTAAAGAGGAGAAATTAACTATGCTACCAGAAGAGTTCCTATGGGGC 3' (SEQ ID NO:51)

3' TTTATTAAGCTTCTCATCAACGGCTATGGTCTTCATTTC 5' (SEQ ID NO:52)

Vector: pQEt; and contains the following restriction enzyme sites 5' Hind III and 3' EcoRI.

OC1/4V endoglucanase (33GP1)

5' AAAAAACAATTGAATTCATTAAAGAGGAGAAATTAACTATGGTAGAAAGACACTTCAGATATGTTCTT 3' (SEQ ID NO:53)

3' TTTTTCGGATCCAATTCTTCATTTACTCTTTGCCTG 5' (SEQ ID NO:54)

Vector: pQEt; and contains the following restriction enzyme sites 5' BamHI and 3' EcoRI.

Thermotoga maritima pullalanase (6GP3)

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5' TTTTGGAATTCATTAAAGAGGAGAAATTAACTATGGAACTGATCATAGAAGGTTAC 3' (SEO ID NO:55)

3' ATAAGAAGCTTTTCACTCTCTGTACAGAACGTACGC 5' (SEQ ID NO:56)

Vector: pQEt; and contains the following restriction enzyme sites 5' EcoRI and 3' Hind III.

The restriction enzyme sites indicated correspond to the restriction enzyme sites on the bacterial expression vector indicated for the respective gene (Qiagen, Inc. Chatsworth, CA). The pQE vector encodes antibiotic resistance (Amp^r), a bacterial origin of replication (ori), an IPTG-regulatable promoter operator (P/O), a ribosome binding site (RBS), a 6-His tag and restriction enzyme sites.

The pQE vector was digested with the restriction enzymes indicated. The amplified sequences were ligated into the respective pQE vector and inserted in frame with the sequence encoding for the RBS. The ligation mixture was then used to transform the $\underline{E.\ coli}$ strain M15/pREP4 (Qiagen, Inc.) by electroporation. M15/pREP4 contains multiple copies of the plasmid pREP4, which expresses the lacI repressor and also confers kanamycin resistance (Kan'). Transformants were identified by their ability to grow on LB plates and ampicillin/kanamycin resistant colonies were selected. Plasmid DNA was isolated and confirmed by restriction analysis. Clones containing the desired constructs were grown overnight (O/N) in liquid culture in LB media supplemented with both Amp (100 ug/ml) and Kan (25 ug/ml). The O/N culture was used to inoculate a large culture at a ratio of 1:100 to 1:250. The cells were grown to an optical density 600 (O.D.600) of between 0.4 and IPTG ("Isopropyl-B-D-thiogalacto pyranoside") was then added to a final 0.6. concentration of 1 mM. IPTG induces by inactivating the lacI repressor, clearing the P/O leading to increased gene expression. Cells were grown an extra 3 to 4 hours. Cells were then harvested by centrifugation.

The primer sequences set out above may also be employed to isolate the target gene from the deposited material by hybridization techniques described above.

Example 2

Isolation of A Selected Clone From the Deposited genomic clones

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A clone is isolated directly by screening the deposited material using the oligonucleotide primers set forth in Example 1 for the particular gene desired to be isolated. The specific oligonucleotides are synthesized using an Applied Biosystems DNA synthesizer. The oligonucleotides are labeled with ³²P--ATP using T4 polynucleotide kinase and purified according to a standard protocol (Maniatis et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Press, Cold Spring, NY, 1982). The deposited clones in the pBluescript vectors may be employed to transform bacterial hosts which are then plated on 1.5% agar plates to the density of 20,000-50,000 pfu/150 mm plate. These plates are screened using Nylon membranes according to the standard screening protocol (Stratagene, 1993). Specifically, the Nylon membrane with denatured and fixed DNA is prehybridized in 6 x SSC, 20 mM NaH₂PO₄, 0.4%SDS, 5 x Denhardt's 500 μg/ml denatured, sonicated salmon sperm DNA; and 6 x SSC, 0.1% SDS. After one hour of prehybridization, the membrane is hybridized with hybridization buffer 6xSSC, 20 mM NaH₂PO₄, 0.4%SDS, 500 ug/ml denatured, sonicated salmon sperm DNA with 1x106 cpm/ml 32P-probe overnight at 42°C. The membrane is washed at 45-50°C with washing buffer 6 x SSC, 0.1% SDS for 20-30 minutes dried and exposed to Kodak X-ray film overnight. Positive clones are isolated and purified by secondary and tertiary screening. The purified clone is sequenced to verify its identity to the primer sequence.

Once the clone is isolated, the two oligonucleotide primers corresponding to the gene of interest are used to amplify the gene from the deposited material. A polymerase chain reaction is carried out in 25 μ l of reaction mixture with 0.5 ug of the DNA of the gene of interest. The reaction mixture is 1.5-5 mM MgCl₂, 0.01% (w/v) gelatin, 20 μ M each of dATP, dCTP, dGTP, dTTP, 25 pmol of each primer and 0.25 Unit of Taq

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polymerase. Thirty five cycles of PCR (denaturation at 94°C for 1 min; annealing at 55°C for 1 min; elongation at 72°C for 1 min) are performed with the Perkin-Elmer Cetus automated thermal cycler. The amplified product is analyzed by agarose gel electrophoresis and the DNA band with expected molecular weight is excised and purified. The PCR product is verified to be the gene of interest by subcloning and sequencing the DNA product. The ends of the newly purified genes are nucleotide sequenced to identify full length sequences. Complete sequencing of full length genes is then performed by Exonuclease III digestion or primer walking.

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Example 3

Screening for Galactosidase Activity

Screening procedures for α -galactosidase protein activity may be assayed for as follows:

Substrate plates were provided by a standard plating procedure. Dilute XL1-Blue MRF E coli host of (Stratagene Cloning Systems, La Jolla, CA) to O.D. $_{600}$ = 1.0 with NZY media. In 15 ml tubes, inoculate 200 μ l diluted host cells with phage. Mix gently and incubate tubes at 37 °C for 15 min. Add approximately 3.5 ml LB top agarose (0.7%) containing 1mM IPTG to each tube and pour onto all NYZ plate surface. Allow to cool and incubate at 37 °C overnight. The assay plates are obtained as substrate p-Nitrophenyl α -galactosidase (Sigma) (200 mg/100 ml) (100 mM NaCl, 100 mM Potassium-Phosphate) 1% (w/v) agarose. The plaques are overlayed with nitrocellulose and incubated at 4 °C for 30 minutes whereupon the nitrocellulose is removed and overlayed onto the substrate plates. The substrate plates are then incubated at 70 °C for 20 minutes.

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Example 4

Screening of Clones for Mannanase Activity

A solid phase screening assay was utilized as a primary screening method to test clones for ß-mannanase activity.

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A culture solution of the Y1090-*E. coli* host strain (Stratagene Cloning Systems, La Jolla, CA) was diluted to O.D.₆₀₀=1.0 with NZY media. The amplified library from *Thermotoga maritima* lambda gtl1 library was diluted in SM (phage dilution buffer): 5×10^7 pfu/µl diluted 1:1000 then 1:100 to 5×10^2 pfu/µl. Then 8 µl of phage dilution (5×10^2 pfu/µl) was plated in 200 µl host cells. They were then incubated in 15 ml tubes at 37 °C for 15 minutes.

Approximately 4 ml of molten, LB top agarose (0.7%) at approximately 52 °C was added to each tube and the mixture was poured onto the surface of LB agar plates. The agar plates were then incubated at 37 °C for five hours. The plates were replicated and induced with 10 mM IPTG-soaked Duralon-UVTM nylon membranes (Stratagene Cloning Systems, La Jolla, CA) overnight. The nylon membranes and plates were marked with a needle to keep their orientation and the nylon membranes were then removed and stored at 4 °C.

An Azo-galactomannan overlay was applied to the LB plates containing the lambda plaques. The overlay contains 1% agarose, 50 mM potassium-phosphate buffer pH 7, 0.4% Azocarob-galactomannan. (Megazyme, Australia). The plates were incubated at 72 °C. The Azocarob-galactomannan treated plates were observed after 4 hours then returned to incubation overnight. Putative positives were identified by clearing zones on the Azocarob-galactomannan plates. Two positive clones were observed.

The nylon membranes referred to above, which correspond to the positive clones were retrieved, oriented over the plate and the portions matching the locations of the clearing zones for positive clones were cut out. Phage was eluted from the membrane cut-out portions by soaking the individual portions in 500 μ l SM (phage dilution buffer) and 25 μ l CHCl₃.

Example 5

Screening of Clones for Mannosidase Activity

A solid phase screening assay was utilized as a primary screening method to test clones for \(\beta \)-monnosidase activity.

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A culture solution of the Y1090-*E. coli* host strain (Stratagene Cloning Systems, La Jolla, CA) was diluted to O.D.₆₀₀=1.0 with NZY media. The amplified library from AEPII 1a lambda gtl1 library was diluted in SM (phage dilution buffer): 5×10^7 pfu/µl diluted 1:1000 then 1:100 to 5×10^2 pfu/µl. Then 8 µl of phage dilution $(5 \times 10^2$ pfu/µl) was plated in 200 µl host cells. They were then incubated in 15 ml tubes at 37 °C for 15 minutes.

Approximately 4 ml of molten, LB top agarose (0.7%) at approximately 52 °C was added to each tube and the mixture was poured onto the surface of LB agar plates. The agar plates were then incubated at 37 °C for five hours. The plates were replicated and induced with 10 mM IPTG-soaked Duralon-UVTM nylon membranes (Stratagene Cloning Systems, La Jolla, CA) overnight. The nylon membranes and plates were marked with a needle to keep their orientation and the nylon membranes were then removed and stored at 4 °C.

A p-nitrophenyl-\$\beta\$-D-manno-pyranoside overlay was applied to the LB plates containing the lambda plaques. The overlay contains 1% agarose, 50 mM potassium-phosphate buffer pH 7, 0.4% p-nitrophenyl-\$\beta\$-D-manno-pyranoside. (Megazyme, Australia). The plates were incubated at 72 °C. The p-nitrophenyl-\$\beta\$-D-manno-pyranoside treated plates were observed after 4 hours then returned to incubation overnight. Putative positives were identified by clearing zones on the p-nitrophenyl-\$\beta\$-D-manno-pyranoside plates. Two positive clones were observed.

The nylon membranes referred to above, which correspond to the positive clones were retrieved, oriented over the plate and the portions matching the locations of the clearing zones for positive clones were cut out. Phage was eluted from the membrane cut-out portions by soaking the individual portions in 500 μ l SM (phage dilution buffer) and 25 μ l CHCl₃.

Example 6

Screening for Pullulanase Activity

Screening procedures for pullulanase protein activity may be assayed for as follows:

Substrate plates were provided by a standard plating procedure. Host cells are diluted to $O.D._{600} = 1.0$ with NZY or appropriate media. In 15 ml tubes, inoculate 200 μ l diluted host cells with phage. Mix gently and incubate tubes at 37 °C for 15 min. Add approximately 3.5 ml LB top agarose (0.7%) is added to each tube and the mixture is plated, allowed to cool, and incubated at 37 °C for about 28 hours. Overlays of 4.5 mls of the following substrate are poured:

100 ml total volume

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0.5g	Red Pullulan Red (Megazyme, Australia)
1.0g	Agarose
5ml	Buffer (Tris-HCL pH 7.2 @ 75 °C)
2ml	5M NaCl
5ml	CaCl ₂ (100mM)
85ml	dH ₂ O

Plates are cooled at room temperature, and thenm incubated at 75°C for 2 hours. Positives are observed as showing substrate degradation.

Example 7

Screening for Endoglucanase Activity

Screening procedures for endoglucanase protein activity may be assayed for as follows:

1. The gene library is plated onto 6 LB/GelRite/0.1% CMC/NZY agar plates (~4,800 plaque forming units/plate) in E.coli host with LB agarose as top agarose. The plates are incubated at 37°C overnight.

- 2. Plates are chilled at 4°C for one hour.
- 3. The plates are overlayed with Duralon membranes (Stratagene) at room temperature for one hour and the membranes are oriented and lifted off the plates and stored at 4°C.
- 4. The top agarose layer is removed and plates are incubated at 37°C for -3 hours.
 - 5. The plate surface is rinsed with NaCl.

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- 6. The plate is stained with 0.1% Congo Red for 15 minutes.
- 7. The plate is destained with 1M NaCl.
- 8. The putative positives identified on plate are isolated from the Duralon membrane (positives are identified by clearing zones around clones). The phage is eluted from the membrane by incubating in $500\mu l~SM + 25\mu l~CHCl_3$ to elute.
- 9. Insert DNA is subcloned into any appropriate cloning vector and subclones are reassayed for CMCase activity using the following protocol:
- i) Spin 1ml overnight miniprep of clone at maximum speed for 3 minutes.
- ii) Decant the supernatant and use it to fill "wells" that have been made in an LB/GelRite/0.1% CMC plate.
 - iii) Incubate at 37°C for 2 hours.
 - iv) Stain with 0.1% Congo Red for 15 minutes.
 - v) Destain with 1M NaCl for 15 minutes.
 - vi) Identify positives by clearing zone around clone.

Numerous modifications and variations of the present invention are possible in light of the above teachings and, therefore, within the scope of the appended claims, the invention may be practiced otherwise than as particularly described.

WHAT IS CLAIMED IS:

- 1. An isolated polynucleotide selected from the group consisting of:
 - (a) SEQ ID NOS: 1-14 and 57-60;
 - (b) SEQ ID NOS: 1-14 and 57-60, wherein T can also be U;
 - (c) polynucleotide sequences complementary to SEQ ID NOS: 1-14 and 57-60;
 - (d) polynucleotide sequences which encode an amino acid sequence as set forth in SEQ ID NOS:15-28, and 61-64; and
 - (e) fragments of (a), (b), (c) or (d) that are at least 15 consecutive bases in length and that will selectively hybridize to DNA which encodes a polypeptide of SEQ ID NOS:15-28, and 61-64.
- 2. A vector comprising a polynucleotide of claim 1.
- 3. A host cell containing the vector of claim 2.
- 4. The method of claim 3, wherein the host cell is a eukaryotic cell.
- 5. The method of claim 3, wherein the host cell is a prokaryotic cell.
- 6. A method for producing a polypeptide comprising:
 - (a) culturing the host cells of claim 3;
 - (b) expressing from the host cell of claim 3 a polypeptide encoded by said polynucleotide; and
 - (c) isolating the polypeptide.

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7. An enzyme selected from the group consisting of:

- (a) an enzyme comprising an amino acid sequence set forth in SEQ ID NOS: 15-28 or 61-64; and
- (b) an enzyme which comprises at least 30 consecutive amino acid residul as an enzyme of (a).
- 8. An enzyme of which at least a portion is coded for by a polynucleotide of claim 1, and which is selected from the group consisting of:
 - (a) an enzyme comprising an amino acid sequence which is at least 70% identical to an amino acid sequence selected from the group of amino acid sequences set forth in SEQ ID NOS:15-28 or 61-64; and
 - (b) an enzyme which comprises at least 30 amino acid residues to the enzyme of (a).
- 9. A method for generating glucose from soluble cell oligosaccharides comprising contacting a sample containing oligosaccharides with an effective amount of an enyzme selected from the group consisting of an enzyme having the amino acid sequence set forth in SEQ ID NOS: 15-28, 61-63 and 64 such that glucose is produced.
- 10. The method of cliam 9, wherein the sample is selected from the group consisting of dairy products, fruit juices, detergents, textiles, guar gum, animal feed, plant biomass and waste products.
- The method of claim 9, wherein the oligosaccharide is selected from the group consisting of maltose, cellobiose, lactose, sucrose, raffinose, stachyose, verbascose, cellulose, starch, amylose, glycogen, disacharrides, polysacharrides and pullulan.

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Figure 1b(Continued)

OC1/4 GLYCOSIDASE - 33G/8 COMPLETE GENE SEQUENCE - 9/95

GENE SEQUENCE - 9/95
ATTE ATTA AGA AGG TOO GAT TITT O'CA AAA GAT TIT ATC TTO GGA ACG GCT ACG GCA TAC 60
THE ARE ARE SET AND Pho Pro Lys AND Phe Ile Pho Clu The ACT GCA GCA TAC 60
61 CAC ATT CAR COM -
61 CAG ATT GAA GGT GCA GCA AAC GAA GAT GGC AGA GGG CCA TCA ATT TGG GAT GTC TTT TCA 120 121 Gln Ile Glu Gly Ala Ala Asn Glu Amp Gly Arg Gly Pro Ser Ile Trp Amp Val Phe Ser 40
and Gly Ale Ale Asn Glu Asp Gly Arg Gly Pro See ATT TGG GAT GTC TTT TCA 120
121 CAC ACC com ene
41 His The Pro Gly Lys The Leu Arn Gly Asp The Gly Asp Val Ala Cys Asp His Tyr His 60
The Leu Ann Gly Asp The Gly Ann Val Ale COT TOT GAC CAT TAT CAC 180
181 CGA TAC AND COLUMN TO THE TOTAL TO THE TYPE HE SO
181 CGA TAC AAG GAA GAT ATC CAG CTG ATG AAA GAA ATA GGG TTA GAC CCT TAC AGG TTC TCT 240 181 CGA TAC AAG GAA GAT ATC CAG CTG ATG AAA GAA ATA GGG TTA GAC CCT TAC AGG TTC TCT 240 241 ATC TCC TGG CCC AGA ATT ACC COMPANY ATT ACC COMPANY AND ALL TAC
ath Leu Het Lys Glu Ile Gly Leu Asp Ala TV Ath TCT TCT 240
241 ATC TCC TGG CCC AGA ATT ATG CCA CIT COC
241 ATC TCC TGG CCC AGA ATT ATG CCA GAT GGG AAG AAC ATC AAC CAA AAG GGT GTG GAT TTC 300 81 Ile Ser Trp Pro Arg Ile Het Pro Asp Gly Lys Asn Ile Asn Gln Lys Gly Val Asp Phe 100 301 TAC AAC AGA CTC GTT CAT CAG
301 The Ash Clark True Ash Clark True 300
101 TAC AAC AGA CTC GTT GAT GAG CTT TTG AAG AAT CAR AND CAR AGA CTC GTT GAT GAG CTT TTG AAG AAT CAR AGA CTC GTT GAT GAG CTT TTG AAG AAT CAR AGA CTC GTT GAT GAG CTT TTG AAG AAT CAR AGA CTC GTT GAT GAG CTT TTG AAG AAT CAR AGA CTC GTT GAT GAG CTT TTG AAG AAT CAR AGA CTC GTT GAT GAG CTT TTG AAG AAT CAR AGA CTC GTT GAT GAG CTT GAT GAT GAG CTT GAT GAG CTT GAT GAT GAG CTT GAT GAT GAT GAT GAT GAT GAT GAT GA
301 TAC AAC AGA CTC GTT GAT GAG CTT TTG AAG AAT GAT ATC ATA CCA TTC GTA ACA CTC TAT 360 101 Tyr Asn Arg Leu Val Asp Glu Leu Leu Lys Asn Asp Ile Ile Pro Phe Val Thr Leu Tyr 120 161 CAC TGG GAC TTA CCC TAG GAG.
361 CAC TCC CAC man and 120
161 CAC TGG GAC TTA CCC TAC GCA CTT TAT GAA AAA GGT GGA TGG CTT AAC CCA GAT ATA GCG 420
121 His Trp Asp Leu Pro Tyr Ala Leu Tyr Glu Lys Gly Gly Trp Leu Asn Pro Asp Ile Ala 140
421 CTC TAT TTC ACA COLUMN 140
421 CTC TAT TTC AGA GCA TAC GCA ACG TTT ATG TTC AAC GAA CTC GGT GAT CGT GTG AAA CAT 480
141 Leu Tyr Phe Arg Ala Tyr Ala Thr Phe Met Phe Asn Glu Leu Gly Asp Arg Val Lys His 160
481 TGG ATT ACA CTG AAC GAA CCA TGG TGT TCT TCT TTC TCG GGT TAT TAC ACG GGA GAG CAT 540 161 Trp Ile Thr Leu Asn Glu Pro Trp Cys Ser Ser Phe Ser Gly Tyr Tyr Thr Gly Glu His 180 541 GCC CCG GGT CAR CAR CAR AND TRY TYP THR Gly Glu His 180
541 GCC CCG GGT CAT CAA AAT TTA CAA GAA GCG ATA ATC GCG GCG CAC AAC CTC TTG AGG GAA 600 601 CAT GGA CAT CCC CTC GTG GU Ala Ila Ila Ala Ala Ala His Asn Leu Leu Arg Glu 200
601 CAT GGA COM
601 CAT GGA CAT GCC GTC CAG GCG TCC AGA GAA GAA GTA AAA GAT GGG GAA GTT GGC TTA ACC 660
201 His Gly His Ala Val Gln Ala Ser Arg Glu Glu Val Lys Asp Gly Glu Val Gly Leu Thr 220 661 AAC GTT GTG ATG ALL SER ARG GLU GLU VAL LYS Asp Gly Glu Val Gly Leu Thr 220
661 AAC GTT GTG ATT ALL 220
ASI VAI VAI VAI ANG TITO COMP NAME OF COLUMN ASI AND COLUMN ASI AND COLUMN ASI VAI VAI AND TO COLUMN ASI VAI VAI AND TO COLUMN ASI VAI VAI AND TO COLUMN ASI
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721 CTT GTT GAT AAG TTC GTT AAT GCA TGG TCC CAT GAC CCT GTT GTT TTC GGA AAA TAT CCC 780 781 GAA GAA GCA CCT GTT GTT AT GCA TGG TCC CAT GAC CCT GTT GTT TTC GGA AAA TAT CCC 780 781 GAA GAA GCA CCT GTT GTT GTT GTT GTT GTT GTT GTT GTT
781 GAA GAA GCA GTT GCA CTT TAT ACG GAA AAA GGG TTG CAA GTT CTC GAT AGC GAT ATG AAT 840
261 Glu Glu Ala Val Ala Leu Tyr Thr Glu Lys Gly Leu Gln Val Leu Asp Ser Asp Het Asn 280
281 Ile Ile Ser Thr Pro Ile Asp Phe Phe Gly Val Asn Tyr Tyr Thr Arg Thr Leu Val Val 300
901 TTT GAT ATG AAC AAT CCT CTT GGA TTT TCG TAT GTT CAG GGA GAC CTT CCC AAA ACG GAG 960
J21 Met Gly Trp Glu Ile Tyr Pro Cle Cla GA TTA TTT GAT ATG CTG GTC TAT CTG AAG GAA ACC
121 Het Gly Trp Glu ile Tyr Pro Gin Gly Leu Phe Asp Het Leu Val Tyr Leu Lys Glu Arg 340
1021 TAT AAA CTA CCA CTT TAT ATC ACA GAG AAC GGG ATG GCT GGA CCT GAT AAA TTG GAA AAC 1080
341 Tyr Lys Leu Pro Leu Tyr Ile Thr Glu Ash Gly Het Ala Gly Pro Asp Lys Leu Glu Ash 160 1081 GGA AGA CTT CAT CAT CAT CAT CAT CAT CAT CAT CA
1081 GGA aca come can all the state of the s
1081 GGA AGA CTT CAT CAT AAT TAC CGA ATT GAA TAT TTG GAA AAG CAC TTT GAA AAA CCA CTT 1140
ory Arg Val His Asp Asn Tyr Arg Ile Glu Tyr Len Glu Lyr AAG CAC TTT GAA AAA GCA CTT 1140
181 Glu Ala Ile Asn Ala Asp Val Asp Leu Lys Gly Tyr Phe Ile Trp Ser Leu Het Asp Asn 400
421 Pro Lys Arg He Leu Lys Asp Ser Ala Het Trp Leu Lys Glu Phe Leu Lys Ser End 419
1317 CTA AAA TCT TAA 1317 CTA AAA TCT TAA 1317 CTA AAA TCT TAA 1317
ye ser and div

STAPHYLOTHERMUS MARINUS GLYCOSIDASE - 12G COMPLETE GENE SEQUENCE 9/95

1 TTG ATA ACC
1 TTG ATA AGG TTT CCT GAT TAT TTC TTG TTT GGA ACA GGT AGA TCA TCG CAC CAG ATC GAG. 60 1 Met 11e Arg Phe Pro Asp Tyr Phe Leu Phe Gly Thr Ala Thr Ser Ser His Gln 11e Glu. 20 61 GGT AAT AAC ATA TTC LICE TTG
61 CCT AAT AAC CO
61 GGT AAT AAC ATA TIT AAT GAT TGG TGG GAG TGG GAG ACT AAA GGG AGG ATT AAG GTG AGA 120
21 Gly Asn Asn Ile Phe Asn Asp Trp Trp Glu Trp Glu Thr Lys Gly Arg Ile Lys Val Arg 40
121 TCG GGT AAG GCA TGT AAT CAT TCG CAA GTG TGT
121 TCG CGT AAG GCA TCT AAT CAT TCG GAA CTC TAT AAA GAA GAC ATA GAG CTT ATC GCT CAG 180 181 CTG CGA TAT AAT CCT TAT AAA CAA GAU ASP Ile Glu Leu Het Ala Glu 60
181 CTG GCA Thm and GTu 60
181 CTG GGA TAT AAT CCT TAT AGG TTC TCC ATA GAG TGG AGT AGA ATA TTT CCC AGA AA. GAT 240 61 Leu Gly Tyr Asn Ala Tyr Arg Phe Ser Ile Glu Trp Ser Arg Ile Phe Pro Arg Lys Asp 80 241 CAT ATA GAT TAT CAG TGG
241 CAT ATA CAT THE CA
241 CAT ATA GAT TAT GAG TCG CTT AAT AAG TAT AAG GAA ATA GTT AAT CTA CTT AGA AAA TAC 300
81 His Ile Asp Tyr Glu Ser Leu Asn Lys Tyr Lys Glu Ile Val Asn Leu Leu Arg Lys Tyr 100
301 GGG ATA GAA CCT GTA ATC ACT CTT CAC CAC TTC ACA AAC CCG CAA TGG TTT ATG AAA ATT 360
101 Gly Ile Glu Pro Val Ile Thr Leu His His Phe Thr Asn Pro Gln Trp Phe Het Lys Ile 120
121 Gly Gly Trp Thr Arg Glu Glu Asn Ile Lys Tyr Phe Ile Lys Tyr Val Glu Leu Ile Ala 140
421 TCC GAG ATTA ANA GROUP TO THE ANA GR
421 TCC GAG ATA AAA GAC GTG AAA ATA TGG ATC ACT ATT AAT GAA CCA ATA ATA TAT GTT TTA 480 481 CAA GGA TAT ATT ATT ATT ATT GTT TTA 160 481 CAA GGA TAT ATT ATT ATT ATT ATT GTT TTA 160
481 CAA GGA TAT ATT TCC GGC GAA TGG CCA CCT GGA ATT AAA AAT TTA AAA ATA GCT GAT CAA 540
= **** 944 OGC ATA (COT 111 116
THE CALL AND THE CAT AND COME OF THE CALL AND COME
221 Ile Asn Ile Tyr His Lys Val Asp Lys Ala Phe Asn Trp Gly Phe Leu Asn Gly Ile Leu 240
721 AGG GGA GAA CTA GAA ACT CTC CGT CGA AAA TAC CGA GTT GAG CCC GGA AAT ATT GAT TTC 780 241 Arg Gly Glu Leu Glu Thr Leu Arg Gly Lys Tyr Arg Val Glu Pro Gly Asn Ile Asp Phe 260
781 ATA GGC ATA ANG THE
781 ATA GGC ATA AAC TAT TAT TCA TCA TAT ATT GTA AAA TAT ACT TGG AAT CCT TTT AAA CTA 840 261 Ile Gly Ile Asn Tyr Tyr Ser Ser Tyr Ile Val Lys Tyr Thr Trp Asn Pro Phe Lys Leu 280
"/" '/' LILE TID Ach Dec ob
841 CAT ATT AAA GTC GAA CCA TTA GAT ACA GGT CTA TGG ACA ACT ATG GGT TAC TGC ATA TAT 900
TO AUA LAIR ATT THE CO. A.
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THE SAME LANGE THE TERM IN THE SAME THE
1021 CAC TTA CAA TAC TTA TAT AAA CCC ATG AAT GAA GGA GCA AAG GTG AAA GGA TAT TTC TAC 1080 1081 TCC 1080 TCC 108
1081 TGG AGC TTC ATC CAT AND THE COLUMN TO THE TYPE 360
1081 TGG AGC TTC ATG GAT AAT TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1081 TGG AGC TTC ATG GAT AAT TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1141 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1141 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1141 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1141 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1141 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1141 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1141 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1142 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1144 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1145 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1146 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1147 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1148 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1148 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1148 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1148 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1148 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA CTA GTA 1140 1148 CAA GTT TA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA TTT AAC CAA AGG TTC GGA TTA GTA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA TTT GAG TGG GAT AAA GGA TTT AAC CAA AGG TTC GGA TTT GAG TGG GAT AAA GGA TTT GAG TGG GAT GAT
1141 GAA GTT GAT TAT AAG ACT TTT GAG AGA AAA CCT AGA AAA AGC GCA TAT GTA TAT AGT CAA 1200
AND THE CAR COM AND
OTA OUR CUT ACC ARC ACT ATT ATT
The Lys fyr Gly Leu Lys Asn Leu 470
431 A
421 Glu End 422

Figure 3

Thermococcus 9N1 Glydosidase -318/G Complete gene sequence 9/95

complete game sequence 9/95	
ATG CTA CCA GAA GGC TIT CTC TGG GGC GTG TCC CAG TCC GGC TTT CAG TTC GAG ATG G	
1 Met Leu Pro Glu Gly Phe Leu Trp Gly Val Ser Gln Sex Gly Phe Gln Phe Glu Net G	CC 60
The way was a	
TA CAC AND	
AMP LYS Leu Arg Arg Am Ile Amp Fro Am Thr Am Tro Las TG GTC AGG GAT C	CC 120
21 Amp Lym Leu Ard Arg Amn Ilm Amp Fro Amn The Amp Trp Trp Lym Trp Val Arg Amp P	ro 40
Al Phe Am Ile Lye Arp Glu Leu Val Ser Gly Asp Leu Pro Glu Glu Gly Ile Asp Asp T	180
181 GAA CTT TAC CIC AND	r 60
181 GAA CTT TAC GAG ANG GAT CAC CGC CTC GCC AGA GAC CTC GGT CTG AAC GTT TAC AGG AT	
61 Glu Leu Tyr Glu Lys Asp Kim Arg Leu Ala Arg Amp Leu Gly Leu Am Val Tyr Arg Il	T 240
241 GGA ATA GAG TEG AGG AGG ATC TIT CCC TEG CCA ACG TEG TIT GTG GAG GTT GAC OTT GA 81 Gly Ile Glu Tep ser Arg Ile Phe Pro Tep Pro Thr Tep She Wal Glu Val Asp Val Glu	
101 Arg Asp Ser Tyr Gly Leu Val Lys Asp Val Lys Ile Asp Lys Asp Thr Leu Glu Glu Leu 16) GAG CAG CAG ARA GAG CAG	. 365
The Law Clarks	
THE WAY AND ALL GITS 11T ALM ALM	
131 Amp Glu Ile Ala Am Him Gln Glu Ile Ale Tyr Tyr Arg Arg Vel Ile Glu Him Leu Arg	420
THE TAX OF MAN AND VALUE OF THE PARTY OF THE	140
141 Glu Leu Gly Phe Lys Val Ile Val Asn Leu Asn His Phe The Leu Pro Leu Trp Leu His	480
481 CAT COS 183 ASS.	160
481 GAT CCC ATA ATC CCC ACG CAG CAG CCC CTC ACC CAC GCT AGG ATT GCC TGG CTC GGG CAG 161 Amp Pro Ile Ile Ale Arg Clu Lym Ale Leu Thr Ann Clu Ann The Coc TGG CTC GGG CAG	
161 Amp Pro Ile Ile Ala Arg Clu Lys Ala Leu Thr Aun Cly Arg Ile Cly Trp Val Cly Cln	540
541 GAG ACC CTC CTC CTC CTC CTC CTC	180
541 GAG AGC GTG GTG GAG TTC GGC AAG TAC GGG GGG TAC ATC GGG AAC GCA CTC GGG GAC CTC	444
THE ALE AST ALL CAN GIV AND THE	600
JOI CAT INT THE LOCALOR	200
201 Val ASD Nec Trp Ser Thr Phe Ash Slu Pro Het Val Val Glu Leu Gly Tyr Leu Ala	660
The same was the country of the coun	220
VVA CCC TAC TOT GOT trop con and a	•
221 Pro Tyr Ser Gly Phe Pro Pro Gly Val Met Are Pro Glu Ala Ala Lys Leu Ala Ile Leu	720
	240
144 AAC ATU ATA AAC GOC COR COR COR COR	
241 Asn Met Tie Asn Als His Als Leu Als Tyr Lys Met Ile Lys Lys Phe Asp Arg Val Lys	780
781 COS COS LAS AND	260
781 OCC GAT ANG GAT TOO CGC TOO GAG GCC GAG GTC GGG ATA ATC TAC ANG ANG ANG ATA GCC GCT. 251 Ala Asp Lys Asp Ser Arg Ser Glu Ala Glu Val Gly Tla Tla ANG ANG ANG ANG ANG GCC GCT.	
251 Ale Asp Lys Asp Ser Arg Ser Glu Ale Glu Val Gly Ile Ile Tyr Asn Asn Ile Gly Val	140
\$41 GCC TAT CC3 TAG CAS	280
841 GCC TAT CCA TAC GAC TCC AAC GAC CCA AAC GAC GTG AAA GCT GCA GAA AAC GAC AAC TAC 281 Als Tyx Pro Tyr Asp Ser Asp Asp Fro Lys Asp Vel Lys Asp Cac GAC GAC AAC TAC	•••
281 Als Tyr Pro Tyr Asp Ser Asn Asp Pro Lys Asp Val Lys Als Als Glu Asn Asp Asn Tyr	300 300
JUL THE CAC AGE GGG CTC TTC TTC TTC	300
301 THE CAE AGE GGG CHE THE THE GAE GEA ATE CAE AAG GGE AAG CHE AAC ATE GAG THE GAE	960
The bys bly Lys Lou Asp Ile Clu Phe Am	320
JOI GGT GAC ACC	
121 Gly Glu The Phe Val Lys Val Arg His Leu Arg Gly Asn Ash Trp Ile Gly Val Asn Tyr	1020
AND TED ILE CIV VAL AND TED ILE CIV VAL AND TED	340
AVEL TAC ACE ACE CEA CITY COMP AND	
1021 TAC ACE ACE CAR GRE ACE TAT TOG CAG COO AAC TTO COE ACE ATA COO CTG ATA TOO 341 Tyr Thr Arg Glu Val Val Arg Tyr Ser Glu Pro Lys Phe Pro Ser Ile Pro Leu Ile Ser	1080
The second secon	360
AVOI TTE CCC CCA CTT CAC AND THE THE	
	1140
1141 ACC CCC CTD ACC CAT ACC CAT ACC CAT	380
181 AGG CCC GTA AGG GAC ATC GGC TGG GAG ATC TAT CCG GAG GGG ATC TAC GAG TCG ATA AGA	200
	1200 100
1201 GAG GCC AAC AAA TAC GCC GTG GCT TO GCT	-50
1301 GAG GCC AAC AAA TAC GGG GTC CGG GTT TAC GTC ACC GAA AAC GGA ATA GCC GAT TCA ACT COL GLU ALA ARD LYE TYT CLY VAL PTG VAL TYC VAL MER CLU ACC GAA ATA GCC GAT TCA ACT	1260
the City Ren City Ila Ala App Ser Thr	120
1161 CAC ACC CTG CGG CCG TAC TAC CTG CGG LATER TO THE COMPANY TO T	
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The second ser his Val Ala Lys Ile Glu Clu Ala Tor Olu	
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Figure 4a

461	CTC CC Leu G1	T TI Y Ph	יבא ש יבא ש	NAC NAC	AGG Arg	TTC Phe	CTA CGC	CTC	TAT	` AAA	GTG	LAT	CTC	ATA	ACC	ıγ z Mα	Glu	LLD	Ala	460
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501	CÁA ATO	Arg	CLU	Lys	TTC Pba	GGA Gly	CTT	GRG Gly	TGA End	15 51	10					ora.	Val	ser	Ly s	500

Figure 4b(Continued)

120

ATG GAA AGG ATC GAT GAA ATT CTC TCT CAG TTA ACT ACA GAG GAA AAG GTG AAG CTC जा Asp Glu lie Leu Ser Gln Leu Thr Thr Glu Gln Lys Met Glu Arg He Lys 1.cu Val GTG GGG GTT GGT CTT CCA GGA CTT TTT GGG AAC CCA CAT TCC AGA GTG GriT GCG CCT Val Gly Val Gly Leu Pro Gly Leu Phe Gly Asa Pro His Ser Arg GIY Ala Als GGA GAA ACA CAT CCC GTT CCA AGA CTT GGA ATT CCT GCG TTT GTC CTG 121 CCA GAT CCT 180 CCC Val Pro Arg Leu Gly He Pro 41 Gly Glo Thr His Pro Ala Pho Λla Asp Cly Pru 60 GCA GGA CTC AGA ATA AAT CCC ACA AGG GAA AAC GAT GAA AAC ACT TAC 181 ACG ACG GCA 240 Ala Gly Leu Arg lic Asn Pro Thr Arg Glu Asn Asp Glu Asn Tyr Туг The Thr TIT CCC GIT GAA ATC ATG CTC GCT TCT ACC TGG AAC AGA GAC CTT CTG GAA GAA CTG GGA 300 Phe Pro Val Glu lic Mei Leu Ala Ser The Top Asn Arg Asp Glu Val Gly 100 AAA GCC ATG GGA GAA GAA GTT AGG GAA TAC GGT GTC GAT GTG CTT CTT CCT GCG ATG 360 Lys Als Mci Gly Glu Glu Val Arg Glu Tyr Gly Vel Asp Val Leu Ala Met 120 AND ATT CAR AGA AND COT CTT TGT GGA AGG ANT TTC GAG TAC TAC TOA GAT CCT GTC Asn Ile His Arg Asn Pro Leu Cys Gly Arg Asn Phe Glu Tyr Tyr Ser 420 A.sp Pro Val. 140 CTT TCC GGT GAA ATG GCT TCA GCC TTT GTC AAG GGA GTT CAA TCT CAA 421 CCC GTG GGA Gly Glu Met Ala Ser Ala Phe Val Lys Gly Val Gln Ser GIY TGC ATA AAA CAC TIT GTC GCG AAC AAG CAG GAA ACG AAC AGG ATG GTA CTG GAC ACG ATC 540 161 Lyx His Phe Val Ala Asn Asn Gin Giu The Asn Arg Met Val معہ Thr 180 GTG TCC GAG CGA GCC CTC AGA GAA ATA TAT CTG AAA GGT TTT GAA ATT CCT στc AAG *** 600 Val Ser Glu Arg Ala Leu Arg Glu lle Tyr Leu Lys Gly Phe Glu Ala Val Lys Lys 200 GCA AGA CCC TGG ACC GTG ATG AGC GCT TAC AAC AAA CTG AAT GGA AAA 601 TAC TGT TCA CAG 660 701 Ala Arg Pro Trp Thr Val Met Ser Ala Tyr Asn Lys Leu Asn Gly Lys Tyr 220 Gin Cys Scr ALC GAN TGG CTT TTG ANG ANG GTT CTC AGG GAN GAN TGG GGA TTT GGC GGT TTC 720 CTC ATG Asn Glu Trp Leu Leu Lys Lys Vai Leu Arg Glu Glu Τmp Gly Phe Gly Gly AGC GAC TGG TAC GCG GGA GAC AAC CCT GTA GAA CAG CTC AAG GCC GGA 721 AAC GAT ATG ATC 780 Ser Asp Trp Tyr Ala Gly Asp Asn Pro Vai Clu Gin Leu Lys Afa Gly Asn ۸عې McI He 260 ATG CCT GGG AAA GCG TAT CAG GTG AAC ACA GAA AGA AGA GAT GAA ATA 781 GAA GAA ATC ATG 840 261 GIY LYS Ala Tyr Gin Val Asn Thr Glu Arg Arg Asp Clu Glu Glu 250 lle Met GAG GCG TTG AAG GAG GGA AAA TTG AGT GAG GAG GTT CTC GAT GAG TGT 841 AGA AAC 231 ATT Leu Lys Glu Gly Lys Leu Ser Clu Glu Val Leu Asp Giu Asn Arg CTC AAA GTT CTT GTG AAC GCG CCT TCC TTC AAA GGG TAC AGG TAC TCA AAC AAC CCG GAT 960 301 Leu Lyx Val Lou Val Asn Ala Pro Scr Phe Lys Tyr Arg Tyr Asn 120 CTC GAA TCT CAC GCG GAA GTC GCC TAC GAA GCA GGT GCG GAG GGT GTT 961 321 CTC CIT CIT GAG 1020 Val Ala Tyr Glu Ala Gly Ala Glu Gly Vαl 340 Glu 1021 AAC AAC GOT GTT CTT CCG TTC GAT GAA AAT ACC CAT GTC GCC GTC TTT GGC 1080 Gly Val Leu Pro Phe Asp Glu Asn Thr His Val ACC CGT CAA Cly Thr GIY Gin 360 ATC GAA ACA ATA AAG GGA GGA ACG GGA ACT GGA GAC ACC CAT CCG AGA TAC ACG ATC TCT lle Glu The He 1.9x Gly Gly The Gly See Gly Asp The His 380 Arg Tyr The He Ser HAT ATC CTT GAA GGC ATA AAA GAA AGA AAC ATG AAG ITC GAC GAA GAA CTC Tall lie Leu Giu Giy IIc Lys Giu Acg Ann Mei Lys Phe Anj Giu CCT TCC ۸۲ 1200 Clu The Tyr Sei

Figure:.5a

12DI GAG GAG TAC ATA AAA AAG ATG AGA GAA ACA GAG GAA TAT AAA CCC AGA 401 Glu Glu Tyr He Lyx Lyx Mei Arg Glu Thr Glu Glu Tyr Lyx Pro Arg vc.c. GAC r: T TGG A3₽ Ser Τm 1241 GGA ACG GTC ATA AAA CCG AAA CTC CCA GAG AAT TTC CTC TCA GAA AAA Gly Thr Val lie Lyx Pro Lyx Leu Pro Giu Axa Phe Leu Ser AAG 1320 Ciu Lys He Lys 440 1321 CCT CCA AAG AAA AAC GAT GTT GCA GTT GTG ATC AGT AGG ATC TCC Pro Pro Lys Lvc Asn Asp Val Ala Val Val Val lic CCT GAG GGA TAC Ser Arg He Gly Glu Cly Tyr 460 ISBI GAC AGA AAG CCG GTG AAA GGT GAC TTC TAC CTC TCC GAT GAC GAG CTG 461 Asp Arg Lys Pro Vai Lys Gly Asp Phe Tyr Leu Scr Asp Asp GAA CTC ATA 1440 Glu Leu He Lys 480 1441 ACC GTC TCG AAA GAA TTC CAC GAT CAG GGT AAG AAA GTT GTG GTT CTT Thr Val Ser Lya Glu Phe His Asp Gln Gly Lys Lys Val Val CTG AAC ATC GGA Vai Leu Asn. He Gly ISOL AGT CCC ATC GAA GTC GCA AGC TGG AGA GAC CTT GTG GAT GGA ATT CTT 501 Ser Pro lie Giu Val Ala Ser Trp Arg Asp Leu CIC TGG CAG 1560 Val Asp Gly fle Trp Gin 520 1561 GCG GGA CAG GAG ATG GGA AGA ATA GTG GCC GAT GTT CTT GTG GGA AAG 521 Ala Gly Gin Glu Met Gly Arg Ilc ATT AAT CCC TCC 1620 Val Aia Asp Val Leu Gly Lys Scr 540 1621 GGA AAA CTT CCA ACG ACC TTC CCG AAG GAT TAC TCG GAC GTT CCA TCC City Lys Lou Pro Thr Thr Phe Pro Lys Asp Tyr Scr Asp Val Pro TGG ACG TTC CCA Tπ Thr Pro 1681 GGA GAG CCA AAG GAC AAT CCG CAA AGA GTG GTG TAC GAG GAA GAC ATC 561 Gly Glu Pro Lys Asp Asn Pro Gin Arg Val Val Tyr Glu Glu Asp tle GTG GGA TAC 1740 Tyr Vat. Gly . Tyr 1741 AGG TAC TAC GAC ACC TTC GGT GTG GAA CCT GCC TAC GAA TTC GGC TAC Arg Tyr Tyr Asp Thr Phe Gly Val Glu Pro Ala Tyr Glu Phe Gly Tyr CTC TCT TAC 1800 لحا Tyr 600 1801 ACA AAG TTT GAA TAC AAA GAT TTA AAA ATC GCT ATC GAC GGT GAG ACG The Lys Phe Glu Tyr Lys Asp Leu Lys IIc Ala IIc Asp Gly Glu The CTC ŤCG 1860 Arg 1861 TAC ACG ATC ACA AAC ACT GGG GAC AGA GCT GGA AAG GAA GTC TCA CAG 621 Tyr Thr lie Thr Asn Thr Gly Asp Arg Ala Gly Lya Glu Val CTC TAC ATC 1920 Val Tyr 11c Lys 1921 GCT CCA AMA GGA AMA ATA GAC AMA CCC TTC CAG GAG CTG AMA GCG TTT 641 Ala Pro Lys Gly Lys IIc Asp Lys Pro Phe CYC ** ACA AAA 1980 Gin Giu Leu Lys His Lys Thr Lys 660 1981 CTT TTG AAC CCG GGT GAA TCA GAA GAA ATC TCC TTG GAA ATT CCT CTC Leu Leu Asn Pro Gly Glu Ser Glu Glu lie CTT GCG 2040 Ser Leu Pro Arg 680 Asp Leu Ala 2041 AGT TTC GAT GGG AAA GAA TGG GTT GTC GAG TCA GGA GAA TAC GAG GTC 681 Ser Phe Asp Gly Lys Glu AGG GTC GGT GCA 2100 Trp Vat Val Glu Ser Gly Giu Tyr Clu Gly 2101 TCT TCG AGG GAT ATA AGG TTG AGA GAT ATT TTT CTG GTT GAG GGA GAG Ala 701 Ser Ser Arg Asp lie Arg Leu Arg Asp lie Phe Leu Val Giu AAG AGA TTC 2160 Gly Glu Ĺys Arg 720 2161 CCA TGA 2166 721 Pro End 722

Figure 5b(Continued)

THERMOCOCCUS AEDII12RA GLYCOSIDASE (18B/C) COMPLETE GENE SEQUENCE - 9/95

COMPLETZ GENZ SECURNOS (18B/C)	
COMPLETE GENE SEQUENCE - 9/95 I ATG ATC CAC TGC CCG GTT AAA GGG ATT ATA TCT GAG GCT CCC GGC ATA ACC ATC ACA ATA Het lie His Cys Pro Vel Lys Gly Ile Ile Ser Gly Ala Att GCO ATA ACC ATC ACA ATA	
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61 GAT TTA ACT TO GALLERY	20
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Figure 6

THERMOCOCCUS CHITONOPHAGUS GLYCOSIDASE - 22G COMPLETE SEQUENCE - 9/95

1 TTG CTT CCA CAC AND THE	
1 TTG CTT CCA GAG AAC TTT CTC TGG GGA GTT TCA CAG TCC GGA TTC CAG TTT GAA ATG G 1 Het Leu Pro Glu Asn Phe Leu Trp Gly Val Ser Gln Ser Gly Phe Gln Phe Glu Het G	CC 60
at only the Glu Met C	1 30
61 GAC AGA CTG AGG AGG CAC ATT GAT CCA AAC ACA GAT TGG TGG TAC TGG GTA AGA GAT GAT ASP	M 120
the Map IIP TYP TYP TAI ARD ARD CO	
121 TAT AAT ATC AAA AAA GGA CTA GTA AGT GGG GAT CTT CCC GAA GAC GCT ATA AAT TCA TA	
The sea of	
101 GAA TTA TAT CAC ACA CAC CAC	
The bed Gly Leu Ash The Tor Ash	
441 GGA ATT GAA TGC 100 101	
THE VAL ASP VAL CITY THE CITY	
JUL ATT GAT GAG TOT TAG GGG	
The Ser Lys Asp Ala Leu Clu two	120
161 CTT GAT GAA ATC GCT AAC CAA AGG GAA ATA ATA TAT TAT AGG AAC CTA ATA AAT TCC CTA 121 Leu Asp Glu Ile Ala Asp Gli Arg Glu Ile Ile Top Top Agg AAC CTA ATA AAT TCC CTA	470
the lyr lyr Ary Ash Leu Ile Ash Ser Leu	420 140
421 AGA AAG AGG GGT TTT AAG GTA ATA CTA AAC CTA AAT CAT TTT ACC CTC CCA ATA TGG CTT 141 Arg Lys Arg Gly Phe Lys Val Ile Leu Ash Leu Ash Leu	400
and had san his Phe Thr Leu Pro Ile Tro Leu	480 160
481 CAT GAT CCT ATC GAR TOT ACK GAR AND AND GAR	
and Lys Arg Ash Cly Trp Val Ser	540 180
541 GAA AGG AGT GTT ATA GAG TITT CGS AND THE	
181 Glu Arg Ser Val Ile Glu Phe Ala Lys Phe Ala Ala Tyr Leu Ala Tyr Lys Phe Gly Asp	600 200
601 ATA GTA GAC ATG TGG AGG AGA TOTAL AND	200
201 Ile Val ASP Het Trp Ser Thr Phe Asn Glu Pro Het Val Val Ala Glu Leu Gly Tyr Leu	660
661 GCC CCA TAC TCA GGA TTC GGG GGG GGA GGA GGA GGA GGA GGA GGA GG	220
221 Ala Pro Tyr Ser Gly Phe Pro Pro Gly Val Het Asn Pro Glu Ala Ala Lys Leu Val Met	720
721 CTA CAT ATG ATA AAC CCC CAT CCT TTA CCC	240
721 CTA CAT ATG ATA AAC GCC CAT GCT TTA GCA TAT AGG ATG ATA AAG AAA TTT GAC AGA AAA 241 Leu His Het Ile Asn Ala His Ala Leu Ala Tyr Arg Het Ile Lys Lys Phe Asp Arg Lys	780
781 ANA GCT GAT CCA GAA TCA ANA GAA GCA GAA GCA ANA GCA GAA GCA GAA TCA ANA GC	260
781 AAA GCT GAT CCA GAA TCA AAA GAA CCA GCT GAA ATA GGA ATT ATA TAC AAT AAC ATC GGC 261 Lys Ala Asp Pro Glu Ser Lys Glu Pro Ala Glu Ile Gly Ile Ile Tyr Asn Asn Ile Gly	840
841 GTC ACA TAT GGG TOTAL AND THE STATE OF ASA ASA FILE GLY	280
841 GTC ACA TAT CCG TTT AAT CCG AAA GAC TCA AAG GAT CTA CAA GCA TCC GAT AAT GCC AAT 281 Val Thr Tyr Pro Phe Asn Pro Lys Asp Ser Lys Asp Leu Gln Ala Ser Asp Asn Ala Asn	900
201 THE MED DIE ASP ASP ASP ASP ASP ASP ASP ASP	300
901 TTC TTC CAC AGT GGG CTA TTC TTA ACG GCT ATC CAC AGG GGA AAA TTA AAT ATC GAA TTT 101 Phe Phe His Ser Gly Leu Phe Leu The No. 12 The All AGG GGA AAA TTA AAT ATC GAA TTT	960
The bed in Ala He His Arg Gly Lys Leu Asn Ile Glu Phe	320
961 GAC GGA GAG ACA TTT GTT TAC CTT CCA TAT TTA AAG GGC AAT GAT TGG CTG GGA GTG AAT 321 Asp Gly Glu Thr Phe Val Tyr Leu Pro Tyr Leu (Art Ca)	1020
and the tyr bed bys Gly Ann Amp Trp Leu Gly Val Asn	340
1021 TAT TAT ACA AGA GAA CTC GTT AAA TAC CAA GAT CCC ATG TTT CCA AGT ATC CCT CTC ATA	1080
The Strange Pro Ser Ile Pro Leu Ile	360
1081 AGC TTC AAG GGC GTT CCA CAT TAN GGA TAG	1140
of the dip cys Arg Pro Gly The The Ser Lys Asp	1140 380
1141 CGT AAT CCT GTT AGT GAG ATT CGA TICK GAG GAG	
	1200 400
1201 GTA GCT GCC AAT GAA TAT GGA CTT CCT CTL TIC	
	1260
1261 AAA GAT GTA TTA AGG CCC TAT TAC ATC CCL TOT CAG	420
The state of the s	1320
and the city of the city and th	440

Figure 7a

PCT/US97/22623

WO 98/24799

11/46

1121		AAT	CGT	TAT	GAC	CTO	AGA	CCA	TAC	. 117	CAC	TCG	GCA	TTA	ACC	CAT	٨٨٣	TAC	GAA	TGG	i tao
							•	,	.,.	Jet		Trp	VIE	rea	Thr	Asp	Asn	Tyr	Glu	Trp	460
1381	CCC	TTA	CGG	TTC	ACA	ATC	100														1440
		-							019	₽£0	. , , 1	O10	ANT	A#n	Leu	Ile.	Thr	Lys	Clu	Ara	1440 480
1441 481	LVE	CCC	AGG	AAA	AAG	ACT	GTA	AGA	GTA	TTC	AGA	GAG	ATA	CTT	ATT	AAT	AAT	ccc	CTA	451	1500
										FILE	AL U	010	114	Val	Ile	Asn	Asn	Gly	Leu	Thr	1500 500
1501	AGC	ΥVC	ATC	AGG	**	GAG	ATC	TTA	GAG	CYC	GGG	TAG	15	16							
501	ser	Asn	Ile	Arg	Lys	Glu	Ile	Leu	Clu	Clu	Gly	End	51								

Figure 7b(Continued)

PYROCOCCUS FURIOSUS GLYCOSIDASE - 7G1 COMPLETE GENE SEQUENCE - 10/95

		•						COM	LET	(CORPO	FE 31	QUE	NCI .	. 10/	95						
	1	ATG	TTC	CCT	GAA	480														ATG GO	
	1	Het	Phe	Pro	Clu	T.	110	CIT	LCC	CCT	GTG	GCA	CAX	TCG	CGT	TTT	CAC			ATG GO Het G	
		-	- •••		OIL	rya	Phe	Leu	Trp	Gly	Val	Ala	Gln	Ser	C1.	DF -		1 - 1	GAA	ATG GO	εO
	61	GAT	888	CTC						_				•••	GLY	rne	Cin	Phe	Glu	Het C	Y 20
	2:	Asn		CIC	AGG	AGG	AAT	ATT	GAC	ACT	AAC	ACT	CBT	**	T						
		, Gp	Lys	ren	Arg	Arg	λsn	Ile	GEA	The	λαπ	Thr	A s n	7.50	100	CAC	TGC	CTA	AGG	Het G	G 120
7	21	101										1 1112	~2₽	ırp	Trp	His	Trp	Val	Ara	Ann	. 120
1	41	ALA	AAT	ATA	GAG .	AAA	GGC	CTC	CTT	ACT	CCS	~ ~ ~							•	Asp AAT TA	3 40
	4.1	Thr	מנא	Il a	Glu .	Lys	Glv	Leu	V = 1	701	~~	CAT	CTT	CCC	CAG .	GAG	GGG	ATT	A A C	AAT TA Aan Ty	_
						-			v = 1	Ser	GLY.	Азр	rea	Pro	Glu i	Glu -	Glv	11.	A	VV1 1V	C 160
1	81	CAC :	CTI	TAT (GAG)	AAG (car i	***											~,,	ASD Ty	- 60
	61	Glu :	Leu '	Tyr (Slu i	.Va	A 4 n 1	L	- \	AIT	GCA /	4GA	AAG I	CTG (GGT (CTT :	AAT (CCT		Asn Ty AGA ATA	
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6	11 (iv i	110	111 7	'r c	· ·		CTA :	TTC (CX :	rcc (CA	ACG 2	ACA 1	ב דד	~~~ ~				AT AGO	
		•			LP 3	WE ,	Crg 1	lle 1	he i	ro :	rp g	ro :	rhe i	hr	ha 7	1	wu (TT (IAT :	AT AGO	300
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10	7	'11 B	~ ~ ~	WY 1	CAI	AT A	AC C	IT A	TA G	AA G	AT G	TA	36 1	T~ .		-				-	
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221	GÇI	CCC	CIA	C TC	r ccc	TT	cc.	T CC					_						, .,	T Leu 5 ATA	220
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																					780
791																					260
261	Lys	Ala	Asr	700	2001	101	3.00	CAC	CCI	. сс	GAZ	GIT	GGT	ATA	ATT	*				GGA Gly	
	•		,	. шуз	, A3b) Set	Lys	G1 է	Pro) XI a	Glu	(Va)	GIV	71.	73-	- TAC	AAL	- AAC	ATI	CCY	840
841																					280
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301	25-	2.0	CAC	TCA	GGG	CTG	TTC	TTC	GAG	CCC	ATA	CNC					AAT TAA		-		
441	File	LU 6	H13	5er	Gly	Leu	Phe	Phe	Glu	312	Tla	44	~~~	. GGA	, AAA	CTI	TAX neA	ATA	GAC	TTT	960
961																					320
																					320
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361	Thr	Ph.	L	~~~	UIT	CAA	CCY	TAT	CCC	TAT	GCC	TGC	AG.	CC+	GC N	100					
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401	VAl	Clu	Al a	His	Lys	TVE	Glv	Val	3	U T	TAC	GTC	ycc	GAG	AAC	GGA	ATA Ile	GCG	GAT	TCA	1260
					., -	- , -	219	441	FID	A9T	Tyr	∨a1	Thr	Glu	Asn	Glv	Ile	Ala	A = -	SAF	120
															-	,		~4 =	Aap	7 57	740

Figure 8a

1261 421		-			. – ,		. , .	.,.		~14	261	413	116	LAS	Met	lle	Clu	Lys	Ala	Phe	1320 440
1321	GAG Glu	CAT Asp	GCG	TAT	GAA Glu	GIT Val	Lys	GGC G1 y	TAC	TTC Phe	C YC	TCC Trp	GCA Ala	TTA Leu	ACT Thr	GAC Asp	AAC Aan	TTC Phe	GAG Glu	TGG Trp	1380
1381 461			•		,		~Ly	F114	GIY	reu	LYE	GIU	Va 1	טנע	Leu	Ile	Thr	Lys	Glu	Ara	1440
1441 481			_		•						~ y	GAG Glu	ATA Ile	GTA Val	V) T	AAT Asn	TAA neA	GGT Gly	GTT Val	ACG Th:	1500 500
1501 501	Lys	DAG Lys	ATT Ile	GAA Glu	G) G) u	GAX Glu	TTG Leu	CTG Leu	AGG AEG	GGA Gly	TGA End	15 51									

Figure 8b(Continued)

Bankia gouldi endoglucanase (370F1)

(3/071)
9 18 27 36 45
5' ATG AGA ATA CGT TTA GGG AGG CTM GGG CTM
Met Arg Ile Arg Leu Ala Thr Leu Ala Leu Cym Ala Ala Leu Ser Pro Val Thr
The Leu Ala Inr Leu Ala Leu Cys Ala Ala Leu Ser Pro Val Thr
63 72
TTT CCA CAT AND CTA AN
TTT GCA GAT AAT GTA ACC GTA CAA ATC GAC GCC GAC GGC GGT AAA AAA CTC ATC
Phe Ala Asp Asn Val Thr Val Glr Ile Asp Ala Asp Gly Cly Lys Lys Leu Ile
• • •
AGC CC) GGG GGG GGG AGC 135 144 153 162
AGC CGA GCC CTT TAC GGC ATG AAT AAC TCC AAC GCA GAA AGC CTT ACC GAT ACT Ser Arg Ala Leu Tyr Gly Met Agn Agn Sar Arg Ala Cal GAT ACC
Ser Arg Ala Leu Tyr Gly Mer Asn Asn Ser Asn Ala Glu Ser Leu Thr Asp Thr
171 180 189 198 207 216
THE CAR CAN CON CON CON CON CON CON CON CON CON CO
Asp Trp Gln Arg Phe Arg Asp Ala Gly Val Arg Met Leu Arg Glu Asn Gly Gly
and the first Arg Giu Asn Gly Gly
225 234 243 252 261 272
AAC AAC AGC ACC AAA TAT AAC TOO CLA TOO CLA
Asn Asn Ser The Lys Tyr Asn Trp Gln Leu His Leu Ser Ser His Pro Asp Trp
The state of the s
279 288 297 306 215
TAC AAC AAT GTC TAC GCC CCC AAC AAC AAC AAC AAC AAC AAC A
Tyr Asn Asn Val Tyr Ala Gly Asn her had Tog GAC AAC CGG GTA GCC CTG ATT
Tyr Asn Asn Val Tyr Ala Gly Asn Asn Asn Trp Asp Asn Arg Val Ala Leu Ile
333 342
CAG GAA AAC CTG CCC CCC CCC CCC CCC CCC CCC CCC CC
CAG GAA AAC CTG CCC GGC GCC GAC ACC ATC TGG GCA TTC CAG CTC ATC GGT AAG
Gln Glu Asn Leu Pro Gly Ala Asp Thr Met Trp Ala Phe Gln Leu Ile Gly Lys
387 306
GTC GCG GCG ACT TCT GCC TAG 336 414 423 432
GTC GCG GCG ACT TCT GCC TAC AAC TIT AAC GAT TGG GAA TTC AAC CAG TCG CAA
Val Ala Ala Thr Ser Ala Tyr Asn Phe Asn Asp Try Glu Phe Asn Gln Ser Gln
441 450
TGG TGG ACC GGC GTC GCT CAG AAT CTC GCT GGC GGC GGT GAA CCC AAT CTG GAC
Trp Trp Thr Gly Val Ala Gln Asn Leu Ala Gly Gly Gly Glu Pro Asn Leu Asp
40.5
495 504 513 522 531 540
GGC GGC GGA GCG CTG GTT GAA GGA GAC CCC AAT CTC TAC CTC ATG GAT TGG
Gly Gly Glu Ala Leu Val Glu Gly Asp Pro Asa Leu Tyr Leu Het Asp Trp
549 558 567 576 585 594
TCG CCA GCC GAC ACT GTG GGT ATT CTC GAC CAC TGG TTT GGC GTA AAC GCG CTG Ser Pro Ala Asp Thr Val Gly Tle Leu Asp Via TTT GGC GTA AAC GCG CTG
Ser Pro Ala Asp Thr Val Gly Ile Leu Asp His Trp Phe Gly Val Asn Gly Leu
603 612 621 630 639 648
OCC GIG CGG CGC XXX CCC XXX mac man
Gly Val Arg Arg Gly Lys Ala Lys Tyr Trp Ser Met Asp Asn Glu Pro Gly Ile
For Not hap han Glu Pro Gly Ile
657 666 675 684 693 700
TGG GTT GGC ACC CAC CAR CON
THE CALL CALL CALL CALL CALL CALL
TGG GTT GGC ACC CAC GAC GAT GTA GTG AAA GAA CAA ACG CCG GTA GAA GAT TTC Trp Val Gly Thr His Asp Asp Val Val Log Cly
Trp Val Gly Thr His Asp Asp Val Val Lys Glu Gln Thr Pro Val Glu Asp Phe

Figure 9a

Bankia gouldi endoglucanese (37021) (continued)

720 729 CTG CAC ACC TAT TTC GAA ACC GCC AAA AAA GCC CGC GCC AAA TTT CCC GGT ATT 738 Leu His Thr Tyr Phe Glu Thr Ala Lys Lys Ala Arg Ala Lys Phe Pro Cly Ile 783 AAA ATC ACC GGT CCG GTG CCC GCT AAT GAG TGG CAG TGG TAT GCC TGG GGC GGT 792 Lys Ile Thr Gly Pro Val Pro Ala Asn Glu Trp Gln Trp Tyr Ala Trp Gly Gly 828 TTC TCG GTA CCC CAG GAA CAA GGG TTT ATG AGC TGG ATG GAG TAT TTC ATC AAG 837 Phe Ser Val Pro Gln Glu Gln Gly Phe Met Ser Trp Met Glu Tyr Phe Ile Lyr 891 CGG GTG TCT GAA GAG CAA CGC GCA AGT GGT GTT CGC CTC CTC GAT GTA CTC GAT 900 Arg Val Scr Glu Glu Gln Arg Ala Ser Gly Val Arg Leu Asp Val Leu Asp 936 945 CTG CAC TAC TAC CCC GGC GCT TAC AAT GCG GAA GAT ATC GTG CAA TTA CAT CGC Leu His Tyr Tyr Pro Gly Ala Tyr Asn Ala Glu Asp Ile Val Gln Leu His Arg 990 999 1008 ACG TTC TTC GAC CGC GAC TTT GTT TCA CTG GAT GCC AAC GGG GTG AAA ATG GTA The Phe Phe Asp Arg Asp Phe Val Ser Leu Asp Ala Asn Gly Val Lim Het Val 1044 1053 GAA GGT GGC TGG GAT GAC AGC ATC AAC AAG GAA TAT ATT TTC GGG CGA GTG AAC 1062 Glu Gly Gly Trp Asp Asp Ser Ile Asn Lys Glu Tyr Ila Phe Gly Arg Val Asn 1107 1116 GAT TGG CTC GAG GAA TAT ATG GGG CCA GAC CAT GGT GTA ACC CTG GGC TTA ACC Asp Trp Leu Glu Glu Tyr Met Gly Pro Asp His Gly Val Thr Leu Gly Leu Thr 1161 GAA ATG TGC GTG CGC AAT GTG AAT CCG ATG ACT ACC GCC ATC TGG TAT GCC TCC 1170 Glu Met Cys Val Arg Asn Val Asn Pro Met Thr Thr Ala Ile Trp Tyr Ala Ser 1206 1215 1224 ATG CTC GGC ACC TTC GCG GAT AAC GGC GTC GAA ATA TTC ACC CCA TGG TGC TGG Met Leu Gly Thr Phe Ala Asp Asn Gly Val Glu Ile Phe Thr Pro Trp Cys Trp 1251 1260 1269 AAC ACC GGA ATG TGG GAA ACA CTC CAC CTC TTC AGC CGC TAC AAC AAA CCT TAT Asn Thr Gly Met Trp Glu Thr Leu His Leu Phe Ser Arg Tyr Asn Lys Pro Tyr 1314 1323 1332 CGG GTC GCC TCC AGC TCC AGT CTT GAA GAG TTT GTC AGC GCC TAC AGC TCC ATT Arg Val Ala Ser Ser Ser Ser Lau Glu Glu Phe Val Ser Ala Tyr Ser Ser Ile 1368 1377 1386 AAC GAA GCA GAA GAC GCC ATG ACG GTA CTT CTG GTG AAT CGT TCC ACT AGC GAG

Asn Glu Ala Glu Asp Ala Met Thr Val Leu Leu Val Asn Arg Ser Thr Ser Glu Figure 9b(Continued)

Bankia gouldi endoglucanase (37GP1) (continued)

ACC CAC ACC GCC ACT GTC GCT ATC GAC GAT TTC CCA CTG GAT GGC CCC TAC CGC Thr His Thr Ala Thr Val Ala Ile Asp Asp Phe Pro Leu Asp Gly Pro Tyr Arg

1467 1476 1485 1494 1503 1512
ACC CTG CGC TTA CAC AAC CTG CCG GGG GAG GAA ACC TTC GTA TCT CAC CGA GAC
Thr Leu Arg Leu His Asn Leu Pro Gly Glu Glu Thr Phe Val Ser His Arg Asp

1521 1530 1539 1548 1557 1566
AAC GCC CTG GAA AAA GGT ACA GTG CGC GCC AGC GAC AAT ACG CTA ACA CTC GAG
Asn Ala Lau Glu Lys Gly Thr Val Arg Ala Ser Asp Asn Thr Val Thr Leu Glu

1575 1584 1593 1602 1611
TTG CCC CCT CTG TCC GTT ACT GCA ATA TTG CTC AAG GCC CGG CCC TAA 3'
Leu Pro Pro Leu Ser Val Thr Ala Ile Leu Leu Lys Ala Arg Pro ***

Figure 9a(Continued)

Thermotoga maritima Alpha-qalactosidade Complete Gene Sequence (LC+3)

5. GTG ATC TGT GTG GAA ATA TITC GGA ANG ACC TTC ACA GAG GGA AGA TTC GTT CTC
Val Ile Cys Val Glu Ile Phe Gly Lys Thr Phe Arg Glu Gly Arg Phe Val Leu
AAA GAG AAA AAC TTC ACA CTT GAG TTC GCG GTG GAG AAG ATA CAC CTT GCC TCC Lys Glu Lys Asn Phe Thr Val Glu Phe Ala Val Glu Lys Ile His Leu Gly Trp
117
ANG ATC TOC GGC AGG GTG ANG GGA AGT CCG GGA AGG CTT GAG GTT CTT CGA ACG
Lys Ile Ser Gly Arg Val Lys Gly Ser Pro Gly Arg Leu Glu Val Leu Arg Thr
ANA GCA CCG GAA AAG GTA CTT GTG AAC AAC TCG CAG TCC TCG GGA CCG TCC AGG
Lys Ala Pro Glu Lys Val Leu Val Asn Asn Trp Gln Ser Trp Gly Pro Cys Arg
GTG GTC GAT GCC TTT TCT TTC AAA CCA GCT GAA ATA GAT CGG AAG TGG AGA TAC
Val Val Asp Ala Phe Ser Phe Lys Pro Pro Clu Ile Asp Pro Asm Trp Ary Tyr
279 288 297 306 315 324 ACC GCT TCG GTG GTC CCC GAT GTA CTT GAA AGG AAC CTC CAG AGC GAC TAT TTC
Thr Ala Ser Val Val Pro Asp Val Lou Glu Ary Asm Leu Gln Ser Asp Tyr Phe
333 342 351 360 369 378 CTG CCT GAA GAA GGG TAC GGG TIT CTG AGT TCG AAA ATC GCA CAT CCT
Val Ala Glu Glu Gly Lys Val Tyr Gly Phe Leu Ser Ser Lys Ile Ala His Pro
387 396 405 414 422
THE THE GET GIG GAA GAT GGG GAA CIT GIG GCA TAC CITC GAA TAT THE GAT GIC
Phe Phe Ala Val Glu Asp Gly Glu Leu Val Ala Tyr Leu Glu Tyr Phe Asp Val
441 450 459 460 477
THE GAL GAC TIT GIT COT CIT GAA CCT CITC GIT GIA CITC GAG GAT CCC AAC
Glu Phe Amp Amp Phe Val Pro Leu Glu Pro Leu Val Val Leu Glu Amp Pro Amn
ACA CCC CITI CTT CTG GAG AAA TAC GCG GAA CTC GTC GGA ATG GAA AAC AAC GCG
The Pro Leu Leu Clu Lys Tyr Ala Clu Leu Val Cly Met Glu Asn Asn Ala
549 558 567 576 585 594 AGA GTT CCA AAA CAC ACA CCC ACT CGA TGG TGG AGG TGG TAG CAT TAG TTC CTT
Arg Val Pro Lys His The Pro The Gly Trp Cyr Ser Trp Tyr Ris Tyr Phe Leu

Figure 10a

Thermotoga maritima Alpha-galactosidane Complete Gene Scquence (2 of 1)

·
GAT CTC ACC TOG GAA GAG ACC CTC AAG AAC CTG AAG CTC OCG AAG AAT TTC CC
Asp Leu Thr Trp Glu Glu Thr Leu Lys Asn Leu Lys Leu Ala Lys Aon Phe Pr
657 666 676
TTC GAG GTC TTC CAG ATA GAC CAC CCC TAC CAA AAG CAC ATA GGT GAC TGG CTC
Phe Glu Val Phe Gln Ile Asp Asp Ala Tyr Glu Lys Asp Ile Gly Asp Trp Leu
711 720 779 770
OTG ACA AGA GGA GAC TIT CCA TCG GTG GAA GAG ATG GCA AAA GTT ATA GCG GAA
Val Thr Arg Gly Asp Phe Pro Ser Val Glu Glu Met Ala Lys Val Ile Ala Glu
765 774 783 789
AAC GOT TTC ATC COG GGC ATA TGG ACC GCC CCG TTC AGT GTT TCT GAA ACC TCC
Asm Gly Phe Ile Pro Gly Ile Trp Thr Ala Pro Phe Ser Val Ser Glu Thr Ser
819 828 837 845
CAT GIA TIC AAC GAA CAT COO GAC TOO GTA GTG AAG GAA AAC GGA GAG COG AAG
Asp Val Phe Asm Glu His Pro Asp Trp Val Val Lys Glu Asm Gly Glu Pro Lys
873 882 891 900 909 918
ATC GCT TAC AGA AAC TOG AAC AAA AAG ATA TAC GCC CTC GAT CTT TCG AAA GAT
Met Ala Tyr Ary Asn Trp Asn Lys Lys Ile Tyr Ala Leu Asp Leu Ser Lys Asp
927 936 945 954 963 972 CAG GTT CTG AAC TGG CTT TTC GAT CTC TTC TCA TCT CTG AGA AAG ATG GGC TAC
Glu Val Leu Asn Trp Leu Phe Asp Leu Phe Ser Ser Leu Arg Lys Met Gly Tyr
981 990 999 1008 1017 1026 AGG TAC TIC AAG ATC GAC TIT CTC TTC GCG GGT GCC GTT CCA GGA GAA AGA AAA
Arg Tyr Phe Lys Ile Asp Phe Leu Phe Ala Gly Ala Val Pro Gly Glu Arg Lys
1035 1044 1053 1062 1071 1080 AMG BAC ATA ACA CCA ATT CAG GCG TTC AGA AAA GGG ATT GAG ACG ATC AGA AAA
Lys Asn Ile Thr Pro Ile Gln Ala Phe Arg Lys Gly Ile Glu Thr Ile Arg Lys
1000
GCG GTG GCA GCA GCA GCA GCA GCA GCA GCA GCA GC
Ala Val Gly Glu Asp Ser Phe Ile Leu Gly Cys Gly Ser Pro Leu Leu Pro Ala
THE COL TOC CA'C COC ATC AGO ATA GGA CCT CA'C ACT CCC CCG TTC TGG GGA
Val Gly Cys Val Asp Cly Met Arg Ile Gly Pro Asp Thr Ala Pro Phe Trp Gly
the first the first transfer of the same and

Figure 10 (Continued)

Thermotoga maritima Alpha-galactosidade Complete Gone Sequence (3.51.7)

1197 1206 1215 1224 1233 1242 GAA CAT ATA GAA GAC AAC CCA CCT CCC CCT GCA ACA TOG CCG CTG AGA AAC CCC
Glu His Ile Glu Asp Asn Gly Ala Pro Ala Ala Arg Trp Ala Leu Arg Asn Ala
1251 1260 1269 1278 1287 1296 ATA ACG ACG TAC TIC ATG CAC GAC ACG TIC TOG CTG AAC GAC CCC GAC TOT CTG
Ile Thr Arg Tyr Phe Met His Asp Arg Phe Trp Leu Asn Asp Pro Asp Cys Leu
ATA CTG AGA GAG GAG AAA ACG GAT CTC ACA CAG AAG GAA AAG GAG CTC TAC TYG
The Lau Arg Glu Glu Lys Thr Asp Leu Thr Gln Lys Glu Lys Glu Leu Tyr Ser
1359 1368 1377 1386 1395 . 1404 TAC ACC TOT OGA CTG CTC GAC AAC ATG ATG ATA GAA AGC GAT GAT CTC TOT CTG
Tyr Thr Cys Cly Val Leu Asp Asn Met Ile Ile Glu Ser Asp Asp Leu Ser Leu
GTC AGA GAT CAT GGA AAA AAG GTT CTG AAA GAA ACG CTG GAA CTG CTG GGT GGA
Val Arg Asp His Gly Lys Lys Val Leu Lys Glu Thr Leu Glu Leu Gly Gly
AGA CCA CGG GTT CAA AAC ATC ATG TCG GAG GAT CTG AGA TAC GAG ATC GTC TCG
Arg Pro Arg Val Gln Asn Ile Met Ser Glu Asp Leu Arg Tyr Glu Ile Val Ser
TOT GGC ACT CTC TCA GGA AAC GTC AAG ATC GTG GTC GAT CTG AAC AGC AGA GAG
Ser Gly Thr Leu Ser Gly Asn Val Lys Ile Val Val App Line Find The Glu
TAC CAC CTG GAA AAA GAA GGA AAG TCC TCC CTG AAA AAA AGA GTC GTC AAA AGA
Tyr His Lau Glu Lys Glu Gly Lys Ser Ser Leu Lys Lys Arg Val Val Lys Arg
1629 1638 1647 1656 1665 GAA GAC GGA AGA AAC TTC TAC TTC TAC GAA GAG GGT GAG AGA GAA TGA 3
Glu Asp Gly Arg Asn Phe Tyr Phe Tyr Glu Glu Gly Glu Arg Glu ***

Figure 10c(Continued)

Thermotoga maritima β-mannanase (6000)

		,	9			18			27			36			45			54
5.	λTG	GGG		GGT	GGC		GAC	TCC	TGG	AGC	ccc	TCA	GTA	TCG	GCG	GAA	TTC	
-													~					
	Met	Gly	Ile	Gly	Gly	Asp	qeA	Ser	Trp	Ser	Pro	Ser	Val	Ser	Ala	Glu	Phe	Leu
			63			72			81			90			99			108
	TTA	TTG	ATC	GTT	GAG	CTC	TCT	TTC	GTT	CTC	TTT	GCA	AGT	CAC	GAG	TIC	CIC	***
	Leu	Leu	Ile	Val	Glu	Leu	Ser	Phe	Val	Leu	Phe	Ala	Ser	Asp	Glu	Phe	Val	Lys
			117			126			135			144			153			162
					λλλ	TIC	CCT	CTG	AAC	CCA	λλλ	GAA	TTC					λGC
						n		7 011		63.2	Tant	Glu	Phe					502
	Val	GIu		GIĀ	гÃз		VTG	Leu		GTÅ				A. 0		110	GIY	
			171			180			189			198		CNC	207	Cum	CENC.	216
	AAC	AAC	TAC	TAC	ATG	CAC	TAC	AAG	AGC	AAC		A1G	ATA					
	λsn	λsn	Tyr	Tyr	Met	His	Tyr	Lys	Ser	nak	G17	Het	11e	ХЕР	Ser	Val	Leu	Glu
			225			234			243			252			261			270
	AGT	GCC	AGA	GAC	λΤΌ	GGT	ATA	λλG	GTC	CTC	λGA	ATC	TGG	CCT	TTC	CTC	GAC	GGG
										 -								
	Ser	Ala	λrg	λsp	Met	Gly	Ile	Lys	Val	Leu	Arg	Ile.	Trp	Gly	Phe	Leu	γsb	Gly
			279			288			297			306			315			324
	GλG	AGT	TAC	TGC	λGλ	GAC	λAG	XXC	YCC	TAC	λTG	CAT	CCT	GAG	ccc	GGT	GIT	TTC
																	17-1	Dhe
	Glu	Ser	Tyr	Cys	Arg		ry#	ASD		тут	nec		Pro	GIU			Val	
			333			342			351	~~		360	TTC	CNN	369		CNC	378
	GGG			GAA														
	Gly												Phe	Glu	λrg	Leu	qeA	Tyr
			387			396			405			414			423			432
	ACA	GIT	GCG	λλλ	GCG	λλλ	GAA	CTC	GGT	λΤλ	λλλ	CTT	GTC	λTT	GTT	CII	GTG	AAC
	Thr	Val	Ala	Lys	Ala	Lys	Glu	Leu	GJA	Ile	Lys	Leu	Val	Ile	Val	Leu	Val	Asn
			441			450			459			468			477			486
	AAC	TGG	GAC	GAC				ХTG	AAC	CAG	TAC	GTG	AGG	TGG	TII	GGA	GGA	ACC
											~			7	Dha			The
	Asn	Trp	yeb) Азр	Phe			Met			. LYI		. Arg	rrp			GIA	
			495			504		~	513		3.00	522		CNO	531		110	540 TAC
								GAT	نکلانا	AAG	ATC				17(TAC
	Ris	His	Asp	yab	Phe	Tyr	Arg	Asp	Glu	Lys	Ile	Lys	Glu	Glu	Ty	Lys	Lys	Tyr

Figure 11a

	T	ber	moto	ga	MAX	itim	La f	-ma	Dhan		(384	100 }-	(c	onti	.Due	a) (6 G P 2	1)
	,	549			55			56	7		576	5		58	5		594	
GTC :	rcc	TT	, CIG	CT.	λλλ	כ כא	T GT	כ אא	T AC	C TAC	ACC	GG	A GT	ר ככי	r ta	C AC	S GAA	
Val s	ser	Phe	: Lei	ı Va.	l As:	n Hi	s Va	l As	n Th	г Туг	The	Gly	/ Val	Pro	Ty:	r Arg	Glu	
		603			61	2		62	1		630)		639	,			
GAG C	CC	ACC	ATC	: ATC	GC	TC	GA	CT	T GC.	A AAC	CAR		CCC	TGT	• GX(: 200	648 CNC	
Glu F	10	III	116	net	- 14	Tr	Gli	ı Leı	ı Ala	nek e	Glu	Pro	Arg	C\2	Glu	The	Asp	
		657			666	;		675	5		684			693			707	
AAA T	CG	GGG	XXC	YCC	CIY	GI1	, CYC	TGC	GTC	3 AAG	GλG	λTG	AGC	TCC	TAC	λτλ	AAG	
Lys S					₩ Q (, vai				Lys	Glu	Met	Ser	Ser	TYI	Ile	Lys	
		711		•	720			729			738			747			756	
AGT C	IG	GAT	CCC	AAC	CAC	CTC	GIG	GCT	GIG	GCG	CYC	GAA	GGA	TTC	TTC	AGC		
										~								
Ser L				A-011	UIB	reu	ATT	ALA	VAI	GIY	Asp	Glu	Glγ	Phe	Phe	Ser	Asn	
		765			774			783			792			801			910	
TAC G	LA (GGλ	TTC	XXX	CCT	TAC	CCT	GGA	GAA	GCC	CNC	TGG	GCC		AAC	GGC		
Tyr GI		ar A	FIIG	PAR	PTO	TYT	GIĀ	GIA	Glu	λla	Glu	Trp	λla	Tyr	Asn	Gly	Trp	
		319			828			837			846			855			061	
TCC GG	T	TT	GAC	TGG	λλG	λλG	CTC	CTT	TCG	ATA	GAG	ACG	CTG	GAC	TTC	CCC	864 ACG	
										~								
Ser Gl	y	41	ASD	Trp	ràs	rys	Leu	Fen	Ser	Ile	Glu	Thr	Val	Хsр	Phe	Gly	Thr	
		173			882			891			900			909			918	
TTC CA	c c	יייי אייי	TAT	CCG	TCC	CAC	TGG	CCI	GTC	AGT	CCA (GAG	***	m= m	GCC			
													_					
Phe Hi			.yr	FIU	241	ura	IIP	CIÀ	VAI	Ser	Pro (Glu	Asn	Tyr	уŢа	Gln	Trp	
		27			936			945			954			963			972	
GGA GC	Gλ	AG 1	rgg .	ATA	GAA	GAC	CAC	ATA	AAG	ATC (GCA A	AAA	GAG	ATC	GGA		-	
										~								
Gly Al		·y ->	·	116	GIG	veb	uiz	116	гЛа	TIG '	NIA I	ГХЗ	Glu	Ile	Gly	ГЛя	Pro	
		81			990			999		1	800		1	017		1	026	
GTT GT	тс	TG	GAA (GAA	TAT	GCX	ATT	CCY	λλG	AGT (GCG (CCA	GTT	AAC .	AGA	ACG (GCC	
Val Va		eu (aru (oru .	IYE	стÃ	116	PTO	Lys	Ser i	NIA :	Pro	Val	Asn .	Arg	Thr .	Ala	
		35			044		1	053		1	062		1	071			000	
ATC TAI	CA	GA (CTC '	rgg .	AAC	GAT	CTG	GTC	TλC	GAT	TC (GGT	GGA	GAT	GGA	ece T	080 ATC	
Ile Ty	ΓÀ	xg 1	Leu 1	rrp .	Asn	Asp	Leu	Val	Tyr	Asp :	Leu (G) A	Gly	Asp	Gly	Ala	Met	

Figure 11b(Continued)

Thermotoga maritima β -mannanase (mag) (continued) (6 6
1089
TTC TCC ATC CTC CCC 1110 111c
Phe Trp Met Leu Ala Gly Ile Gly Glu Gly Ser Asp Arg Asp Glu Arg Gly Tyr
1143 1152
TAT CCC GAC TAG COST 1161 1170 1170
THE AGA ATA GTG AAC GAC AGT CGA CALL
Tyr Pro Asp Tyr Asp Gly Phe Arg Ile Val Asn Asp Asp Ser Pro Glu Ala Glu
CTG ATA AGA GAA TAC GCG AAC GTG 1224 1233
Leu Ile Arg Glu Tyr Ala Lys Leu Phe Asp Thr Gly Gly
Leu Ile Arg Glu Tyr Ala Lys Leu Phe Asn Thr Gly Glu Asp Ile Arg Glu Asp
ACC TCC TCT TTC 1700 1700 1700 1700 1700 1700 1700 170
1296
Thr Cys Ser Phe Ile Leu Pro Lys Asp Gly Met Glu Ile Lys Lys Thr Val Glu
of the City field Lys Lys Thr Val Clu
1.105
1361 1360
Val Arg Ala Gly Val Phe Asp Tyr Ser Asp The Pho Ci
and the fit Lys Leu Ser Val Lys
1459 1970
GTC GAA GAT CTG GTT TTT GAA AAT GAG ATA GAG CAT CTC GGA TAC GGA ATT TAC
Val Glu Asp Leu Val Phe Glu Asn Glu Ile Glu Fig.
Val Glu Asp Leu Val Phe Glu Asn Glu Ile Glu His Leu Gly Tyr Gly Ile Tyr
1411 1200 .
1413 1422 1431 1440 1449 1458 GGC TTT GAT CTC GAC ACA ACC CGG ATC CCG GAT GGA GAA CAT GAA ATG TTC CTT
THE STATE OF THE S
Gly Phe Asp Leu Asp Thr Thr Arg Ile Pro Asp Gly Glu His Glu Met Phe Leu
1467 1486
What later that the contract the second seco
Glu Gly His Phe Gln Gly Lys Thr Val Lys Asp Sep 5
Glu Gly His Phe Gln Gly Lys Thr Val Lys Asp Ser Ile Lys Ala Lys Val Val
and see the Lys Ala Lys Val Val
THE CAN GAR GIT CAT TITL TOO TOTAL COLUMN TO THE COLUMN
Asn Glu Ala Arg Tyr Val Leu Ala Glu Glu Val Arm Transcription
Asn Glu Ala Arg Tyr Val Leu Ala Glu Glu Val Asp Phe Ser Ser Pro Glu Glu
1575 1604
AAA AAC TGG TGG AAC AGC GGA ACC TGG CAG ACC TGG CA
Val Lys Asn Trp Trp Asn Ser Gly Thr Trp Glp All Gl
Val Lys Asn Trp Trp Asn Ser Gly Thr Trp Gln Ala Glu Phe Gly Ser Pro Asp
Gly Ser Pro Asp

Figure 110 (Continued)

		The	Гъо	tog	A 2	ari	tim.	- 1	3- z a	nna	nas	•	Œ.	E H	(cont	inu	đ)	(60	(يې م :
A.	rr G	16 አእ ፕ	29 GG 2	AC	GGT.	1638 Gag	GI	G GG	164 X AJ	17 NT G	GA (I CA	656 CTG	: CA	നേ	16	65 AC G	NG A	1674 UL CTO	,
11	le G	lu T	TP A	 -sn (Gly	Glu	Val	Gl	 γ λs	in G	 ly)	lla	 Leu	Gl	n Le	u As	n Va	1 Ly	's Leu	
		16	83		1	692			120										1728 A CTC	
Pr	o G1	y Ly	s S	er A	\sp	Trp	Glu	Glu	Va.	 l Ar	g V	al J	la	Arg	 Ly	s Ph		 u Ar	 g Leu	
TC	A GA	173 A TG 	T C	\С A	TC (746 CTC	GAG	TAC	1755 GAG	TA :	C T	AC A	64 TT	CCX	AAC	177 GT	3 C GAG	G GZ	1782 CTC	
Se	c Gl	и Су	s G1	u I	le I	ieu I	Glu	Тут	λs	Il	e 13	r I	le	Pro	Asr	 1 Va	l Glu	Gly	Leu	
AA C	GC:		1 3 TT 				IXC	GCG	GIT	CIC	3 XX	.c c	CC (GGC	TGG	1827	λAG	ATA	183 <i>6</i> GGC	
Lys	G13	1845		u Az												Val	Lys	Ile	Gly	
CTC	GAC	ATC		2 AA	C G	54 CG A	AC .	GT.	GAA	AGT	, ec	G GA	KG A	TC	ATC	1881 ACT	TTC	ccc	1890 GGA	
ren		. Met 1899					sn'										Phe	Gly	Gly	
AAA	GAG	TAC	YCY	λG	λ T	rc c	AT (STA .	λGλ	ATT	CY(TT	C G	AC	AGA	1935 1021	GCG	GGG :	GTG	
	:	1953			196	2		1	071				_				λla			
AAA 	GAA	CTT	CAC	λT	λ GG	A G	rr c	TC (971 GGT	Cat	CAT	, CI	O G A	GG	TAC 1	989. GAT	GGA	CCG	.998 ATT	
ГЛя	Glu	Leu	His	Ile	e Gl	y Va	al V	al (Sly	Дзр	His	Lei	u Ai	rg '	ľγr	Asp	Gly	Pro	 Ile	
TTC		2007 GAT			201 3 AG	6 A C1	T T	20 AT 2)25 UA :	λGA	ACA	2034 GC2	4 4 G0	ST 2	2 ATG	043 TGA	י נ			
Phe																	-			

Figure 11d (Continued)

ARPII la β -mannosidase (63GB1

5' ATG CTA CCA GAA CAG 777 36 45
54
Met Leu Pro Glu Glu Phe Leu Trp Gly Val Gly Gln Ser Gly Phe Gln Phe Glu
and the Let Trp Gly Val Gly Gln Ser Gly Phe Gln Phe Glo
63 72 81 90
ATG GGC GAC AAG CTC AGG AGG CAC ATC GAT CCA AAT ACC GAC TGG TGG AAG TGG
Met Gly Asp Lys Lou And Tog
Met Gly Asp Lys Leu Arg Arg His Ile Asp Pro Asn Thr Asp Trp Trp Lys Trp
GTT CGC GAT CCT TTC AAC ATA AAA AAG GAG CTT GTG AGT GGG GAC CTT CCC GAG
VAL AND AND THE STATE OF THE ST
Val Arg Asp Pro Phe Asn Ile Lys Lys Glu Leu Val Ser Gly Asp Leu Pro Glu
GAC GGC ATC AAC AAC TAC GAA CTT TTT GAA AAC GAT CAC AAG CTC GCT AAA GGC
ASD GIV IIA AND AND AND AND AND GOOD AN
Asp Gly Ile Asn Asn Tyr Glu Leu Phe Glu Asn Asp Ris Lys Leu Ala Lys Gly
275
CTT GGA CTC AAC GCA TAC AGG ATT GGA ATA GAG TGG AGC AGA ATC TTT CCC TGG
Leu Cly Lou ATT CCC TGG
Leu Gly Leu Asn Ala Tyr Arg Ile Gly Ile Glu Trp Ser Arg Ile Phe Pro Trp
1/9 300
CCG ACG TGG ACG GTC GAT ACC CAG 970 306 315 324
CCG ACG TGG ACG GTC GAT ACC GAG GTC GAG TTC GAC ACT TAC GGT TTA GTA AAG
Pro Thr Trp Thr Val Asp Thr Glu Val Glu Phe Asp Thr Tyr Gly Leu Val Lys
333 949
GAC GTT AAG ATA GAC AAG TCC ACC CTT GCT GAA CTC GAC AGG CTG GCC AAC AAG
THE AND THE ACC CITY GET GAA CITE GAC AGG CITG GEE AAC AAG
Asp Val Lys Ile Asp Lys Ser Thr Leu Ala Glu Leu Asp Arg Leu Ala Asn Lys
387 396 405 414 423 423
GAG GAG GTA ATG TAC TAC AGG CGC GTT ATT CAG CAT TTG AGG GAG CTC GGC TTC
Glu Glu Val Met Tyr Tyr Arg Arg Val Ile Gln His Leu Arg Glu Leu Gly Phe
tal His Leu Arg Glu Leu Gly Phe
AAG GTC TTV CTT > 10 459 468 477
ANG GTC TTC GTT ANC CTC ANC CNC TTC ACG CTT CCA ATA TGG CTC CNC GAC CCG
Lys Val Phe Val Asp Leu Asp His Dr.
Lys Val Phe Val Asn Leu Asn His Phe Thr Leu Pro Ile Trp Leu His Asp Pro
445 604
ATA GTG GCA AGG GAG AAG GCC CTC ACA AAC GAC AGA ATC GGC TGG GTC TCC CAG
The Val Ala Arg Glu Lyc 11 - 1
Ile Val Ala Arg Glu Lys Ala Leu Thr Asn Asp Arg Ile Gly Trp Val Ser Gln

Figure 12a

AEPII la β-mannosidase (63GB1) (continued)

		54.	9		55	3		e r-									
λG	G ÁC	A GT	r GT:	CAC	יכנ מדד ב	י יינרי	~ AAC	. OC.			576			589	5		594 C GGA
Ar	g Th	r Val	l Val	l Glu	2 Phe	Ala	Lys	Тут	Ala	Ala	Tyz	Ile	Ala	His	·	Lei	. Gly
		603															. uzy
ŒΛ	CTY			: גכי	612 TGC	: : NGC	100	621	110	<i>-</i>	630			639)		648 CTC
Ası) La	ı Val	. Asp	Thr	Trp	Sex	Thr	Phe	λsn	Glu	Pro	Met	Val	Val	Val	Glu	Leu
		657			666			675									
GGC	TAC			ccc			GGA	TTT	ccc	CCG	684 GCA	CITY	3.000	693			702 GCC
Gly	XXX	Leu	Ala	Pro	Tyr	Ser	Gly	Phe	Pro	Pro	Gly	Val	Met	Asn	Pro	Glu	λla
		711															
GCG	AAG	cre	GCG	ATC	CTC	AAC	ATG	ATA	110	CCC	738	~~~	~~	747			756
Ala	Lys	Leu	Ala	Ile	Leu	Asα	Het	Il.	Asn	Ala	His	λla	Leu	λla	Tyr	Lys	Met
•		765			774			783									
ATA	AAG	AGG	TTC	GAC		λAG	λAG	GCC	GAT	GAG	792 GAT	ACC.	330	801	~~~		810
									~~~								
Ila	Lyg	Arg	Phe	ζsp	The	Lys	Lys	λla	מבע	Glu	λsp	Ser	Lys	Ser	Pro	λla	Asp
		819			828			837			846			0 5 5			
GIT	GGC	λΤλ	ATT	TAC	AAC	AAC	A TYC	~~	~700	~~~	D10			855			864
							~	GGI	a L.L.	بابات	TAC	CCT	AAA	GXC	CCT	330	CAT
15- 3								~~~									
Val		Ile						~~~									
Val							Ile	Gly		Ala	Тух		Lys .	Asp		<b>λ</b> 3η	λsp
	Gly	11a 873	lle	ŢYŦ	Asn 882	Asn	Ile	Gly '	Val .	Ala	Tyx	Pro	Lys .	Asp	Pro	 λ <b>s</b> n	λsp
ccc 	Gly	11e 873 GAC	Ile	Tyr AAA	Asn 882 GCA	Asn	Ile	Gly :	Val .	Ala AAC	Tyr 900 TAC	Pro	Lys .	Asp 909 AGC	Pro	<b>ASTI</b>	Asp 918 TTC
ccc 	Gly	11a 873	Ile	Tyr AAA	Asn 882 GCA	Asn	Ile GAA	Gly :	Val .	Ala AAC	Tyr 900 TAC	Pro	Lys .	Asp 909 AGC	Pro	<b>ASTI</b>	Asp 918 TTC
ccc 	Gly	11e 873 GAC	Ile	Tyr AAA Lys	Asn 882 GCA	Asn	GAA .	Gly 891 AAC (	Val .	AAC (	Tyr 900 TAC	Pro	CAC .	Asp 909 AGC	Pro	Asn CTG	Asp 918 TTC  Phe
CCC  Pro	Gly AAG Lys	873 GAC  Asp	Ile GTT  Val	Tyr AAA Lys	Asn 882 GCA  Ala 936	Asn GCC  Ala	GAA .	Gly : 891 AAC (	Val .	AAC (	Tyr  900  TAC  Tyr  954	Pro	CAC .	Asp 909 AGC  Ser	Pro GGA GGIY	Asn CTG	Asp 918 TTC  Phe
CCC  Pro	AAG Lys GAT	873 GAC  Asp 927 GCC	Ile GTT Val	Tyr AAA Lys	Asn 882 GCA Ala 936 AAG	Asn GCC Ala	GAA .	Gly 891 AAC AAR A	Val .	AAC AAC AAAC AAAC AAAC AAAC AAAC AAAC	Tyr 900 TAC Tyr 954 GAG	Pro Pro Phe	CAC .	Asp 909 AGC Ser 963	Pro GGA Gly	ASTI CTG  Leu	Asp 918 TTC Phe 972
CCC  Pro	AAG Lys GAT	873 GAC  Asp	Ile GTT Val	Tyr AAA Lys	Asn 882 GCA Ala 936 AAG	Asn GCC Ala	GAA .	Gly 891 AAC AAR A	Val .	AAC AAC AAAC AAAC AAAC AAAC AAAC AAAC	Tyr 900 TAC Tyr 954 GAG	Pro Pro Phe	CAC .	Asp 909 AGC Ser 963	Pro GGA Gly	ASTI CTG  Leu	Asp 918 TTC Phe 972
CCC  Pro TTT  Phe	Gly AAG Lys GAT Asp	873 GAC  Asp 927 GCC  Ala	Ile GTT Val	Tyr  AAA  Lys  CAC	Asn 882 GCA  Ala 936 AAG 	Asn GCC Ala GGT Gly	GAA	Gly : 891 AAC : Asn : Grant Color Co	Val .	Ala AAC AAC AAAC AAAC AAAC AAAC AAAC AAA	Tyr  900 TAC Tyr  954 GAG Glu	Pro Phe:	CAC	Asp 909 AGC Ser 963 GGC Gly	GGA Gly GAA	Asn CTG Leu AAC	Asp 918 TTC Phe 972 TTT
CCC  Pro TTT  Phe	Gly AAG Lys GAT Asp	873 GAC  Asp 927 GCC  Ala	Ile GTT Val	Tyr  AAA  Lys  CAC	Asn 882 GCA  Ala 936 AAG 	Asn GCC Ala GGT Gly	GAA	Gly : 891 AAC : Asn : Grant Color Co	Val .	Ala AAC AAC AAAC AAAC AAAC AAAC AAAC AAA	Tyr  900 TAC Tyr  954 GAG Glu	Pro Phe:	CAC	Asp 909 AGC Ser 963 GGC Gly	GGA Gly GAA	Asn CTG Leu AAC	Asp 918 TTC Phe 972 TTT
CCC  Pro TTT  Phe	Gly  AAG Lys  GAT Asp	873 GAC  Asp 927 GCC  Ala 981 GTT	Tile  GTT  Val  ATC  Ile	AAA Lys CAC His	Asn 882 GCA Ala 936 AAG Lys 990 CTA	ASD GCC Ala GGT Gly	GAA	Gly 891 AAC ABN 945 CTC Leu 999	VAL .	AAC (AAC (AAC (AAC (AAC (AAC (AAC (AAC	Tyr  900 TAC Tyr  954 GAG Glu  008	Pro Phe:	CAC	Asp 909 AGC Ser 963 GGC Gly	Pro  GGA  Gly  GAA  TAC	ASD CTG Leu AAC ASD	Asp 918 TTC Phe 972 TTT Phe 026
CCC  Pro TTT  Phe	Gly  AAG Lys  GAT Asp	873 GAC  Asp 927 GCC  Ala	Tile  GTT  Val  ATC  Ile	AAA Lys CAC His	Asn 882 GCA Ala 936 AAG Lys 990 CTA	ASD GCC Ala GGT Gly	GAA	Gly 891 AAC ABN 945 CTC Leu 999	VAL .	AAC (AAC (AAC (AAC (AAC (AAC (AAC (AAC	Tyr  900 TAC Tyr  954 GAG Glu  008	Pro Phe:	CAC	Asp 909 AGC Ser 963 GGC Gly	Pro  GGA  Gly  GAA  TAC	ASD CTG Leu AAC ASD	Asp 918 TTC Phe 972 TTT Phe 026
CCC Pro TTT Phe GTA Val	Gly  AAG Lys  GAT Asp  AXA Lys	873 GAC  Asp 927 GCC  Ala 981 GTT  Val	GTT Val  ATC Ile	Tyr  AAA  Lys  CAC  CAC  His	ASN  882 GCA Ala  936 AAG Lys  990 CTA Leu	AAA GCC GCC GCT Gly	GAA	Gly: 891 AAC ( AAS ) 45 CTC / Leu / AAAS )	VAL .	AAAC AAAC AAAAC AAAAAC AAAAC AAAAC AAAAC AAAAAC AAAAAC AAAAAA	900 TAC : Tyr : Tyr : 300 TAC : 300	Pro Pro Pro Pro GGC (	CAC His	Asp 909 AGC Ser Ser GGC GGL GGL GGL AAAC	Pro  GGA  Gly  GAA  TTAC	ASTA CTG Leu AAC ITAC	ASP 918 TTC Phe 972 TTT Phe 026 ACC Thr
CCC Pro TTT Phe GTA Val	Gly AAG Lys GAT Asp	873 GAC  Asp 927 GCC  Ala 981 GTT  Val	GTT Val  ATC Ile	Tyr  AAA  Lys  CAC  CAC  His	ASN  882 GCA Ala  936 AAG Lys  990 CTA Leu	AAA GCC GCC GCT Gly	GAA	Gly: 891 AAC ( AAS ) 45 CTC / Leu / AAAS )	VAL .	AAAC AAAC AAAAC AAAAAC AAAAC AAAAC AAAAC AAAAAC AAAAAC AAAAAA	900 TAC : Tyr : Tyr : 300 TAC : 300	Pro Pro Pro Pro GGC (	CAC His	Asp 909 AGC Ser Ser GGC GGL GGL GGL AAAC	Pro  GGA  Gly  GAA  TTAC	ASTA CTG Leu AAC ITAC	ASP 918 TTC Phe 972 TTT Phe 026 ACC Thr
CCC Pro	Gly  AAG Lys  GAT Asp  AAA Lys	873 GAC  Asp 927 GCC  Ala 981 GTT  Val	Ile GTT Val ATC	AAA Lys CAC CAC His	882 GCA Ala 936 AAG CTA LLYS	ASIN GCCC Alla GGT GIY	GAA	Gly : September 1	Val .	AAAC (AAAC (	900 TAC : TTY : TT	Pro	CAC His Asp	909 AGC Ser Ser GGC GGC AAC	Pro  GGA  GGA  GAA  TAC  TYT  CTC	ASTA  CTG  Leu  AAC  ASTA  1  TYE  ATA	ASP 918 TTC Phe 972 TTT Phe 026 ACC Thr

Figure 12b(Continued)

# AEPII la β-mannosidase (630B1) (continued)

(continued)
TTC AAG GGC GTT CCC AAC TAC GGC TAC TCC TGC AGG CCC GGC ACG ACC TCC GCC  Phe Lys Gly Val Bro Acc TCC TGC AGG CCC GGC ACG ACC TCC GCC
Phe Lys Gly Val Pro Asn Tyr Gly Tyr Ser Cys Arg Pro Gly Thr Thr Ser Ala  1143 1152 1161 1170 1179 1188
GAT GGC ATG CCC GTC AGC GAT ATC GGC TGG GAA GTC TAT CCC CAG GGA ATC TAC Asp Gly Met Pro Val Ser Asp Ile Gly Trp Glu Val Tyr Pro Gln Gly Ile Tyr
GAC TCG ATA GTC GAC GCC ACC AAG TAC AGT GTT CCT GTT TAC GTC ACC ACC AAG TAC AGT GTT CCT GTT TAC GTC ACC ACC ACC ACC ACC ACC ACC ACC ACC A
1251 1260 1269 1278 1287 1296 GGT GTT GCG GAT TCC GCG GAC ACG CTG AGG CCA TAC TAC ATA GTC ACG CTG ACG
1305 1314
1305 1314 1323 1332 1341 1350 TCA AAG ATA GAG GAA GCC ATT GAG AAT GGA TAC CCC GTA AAA GGC TAC ATG TAC Ser Lys Ile Glu Glu Ala Ile Glu Asn Gly Tyr Pro Val Lys Gly Tyr Met Tyr
TGG GCG CTT ACG GAT AAC TAC GAG TGG GCC CTC GGC TTC AGC ATG AGC TTT GCD
1413
CTC TAC AAG GTC GAC CTC ATC TCC AAG GAG AGG ATC CCG AGG GAG AGA AGC GTT Leu Tyr Lys Val Asp Leu Ile Ser Lys Glu Arg Ile Pro Arg Glu Arg Ser Val
GAG ATA TAT CGC AGG ATA GTG CAG TCC AAC GGT GTT CCT AAG GAT ATC AAA GAG
1521 1530
Glu Phe Leu Lys Gly Glu Glu Lys ***

Figure 12C(Continued)

# OCI/4V Endoglucanase (33GP1)

5' ATG GTA GAA AGA CAC TTC AGA TAT GIT CTT ATT TGC ACC CTG TTT CTT GTT A	5.
Met Val Glu Arg His Phe Arg Tyr Val Leu Ile Com	ī
nec Val Glu Arg His Phe Arg Tyr Val Leu Ile Cys Thr Leu Phe Leu Val Mo	et
CTC CTA ATC TCA TCC ACT CAG TGT GGA AAA AAT GAA CCA AAC AAA AGA GTG AA	08
Leu Leu Ile Ser Ser Thr Gln Cys Gly Lys Asn Glu Pro Asn Lys Arg Val As	\T
AGC ATG GAA CAG TCA GTT GCT GAA AGT GAT AGC AAC TCA GCA TTT GAA TAC AAC	
Ser Met Glu Gln Ser Val Ala Glu Ser Asp Ser Asn Ser Ala Phe Glu Tyr Asn	<u>-</u>
1	n
171 180 189 198 207 716	5
AAA ATG GTA GGT AAA GGA GTA AAT ATT GGA AAT GCT TTA GAA GCT CCT TTC GAA	
Lys Met Val Gly Lys Gly Val Asn Ile Gly Asn Ala Leu Glu Ala Pro Phe Glu	•
775	į.
GGA GCT TGG GGA GTA AGA ATT GAG GAT GAA TAT TTT GAG ATA ATA	
Charles and the transfer of th	
Gly Ala Trp Gly Val Arg Ile Glu Asp Glu Tyr Phe Glu Ile Ile Lys Lys Arg	
279 280 227	
GGA TTT GAT TCT GTT AGG ATT CCC ATA AGA TCC TCA CCC 315	
GIV Pho And Garage	
Gly Phe Asp Ser Val Arg Ile Pro Ile Arg Trp Ser Ala His Ile Ser Glu Lys	
333 342 25	
CCA CCA TAT GAT ATT GAC AGG AAT TTC CTC GAA AGA GTT AAC CAT GTT GTC GAT	
Pro Pro Tyr Asp Ile Asp Arg Asp Db. Law Co.	
Pro Pro Tyr Asp Ile Asp Arg Asn Phe Leu Glu Arg Val Asn His Val Val Asp	
387 395	
ANG GCT CTT GAG AAT AAT TTA ACA GTA ATC ATC ARC ACC GTG 432	
Arg Ala Leu Glu Asp Asp Leu The Car Titt GAA GAA	
Arg Ala Leu Glu Asn Asn Leu Thr Val Ile Ile Asn Thr His His Phe Glu Glu	
441 450	
TAT CAN GAN CCG GAT ANA TAC GGC CAT CTT CTT	
Leu Tyr Gln Glu Pro Asp Lys Tyr Gly Asp Wall to	
Leu Tyr Gln Glu Pro Asp Lys Tyr Gly Asp Val Leu Val Glu Ile Trp Arg Gln	
495 504 555	
AND THE THE ANA GAT TAC CCG GAA AAT CTG TTC TTC CALL	
Ile Ala Lys Phe Phe Ive Acc and	
Ile Ala Lys Phe Phe Lys Asp Tyr Pro Glu Asn Leu Phe Phe Glu Ile Tyr Asn	

Figure 13A

			49	C1/	47	<b>E</b> ndo	glu	CADA		(330	3 <b>P</b> 1)	( c	onti	nue	1)		
c	λG C	ירים כ	- m	'AC 1		558			567			576		S	85		5.0
_				.AG /	MC :	rig i	ICA (	GCT (	EXX )	W 1	LCC 1	NAC (	GCA (	TTT	አፓ ር	۲x .	59 WA GT
G	lu P	LO Y	la G	ln /	ısn I	eu 1	hr )	lla G	lu I	ys T	ر طعر د طعر	usn /	la I	eu T	 yr P	ro L	WA GT
C	ፐር ኢ	6 AAG	03 TT A	ምር a	6 6	12	<b>.</b>	6	21		6	30		6.	39		64
-											GG A	TT G	TC A	TT A	LC C	AT G	64: CT CC!
L	eu Ly			le A	rg G	lu S	er A	sn P	ro T	hr A	rg I	le V	al I	le II	le As	ъ A.	 la Pro
J.	C TO	6: 3G G(	57 DAC	AC TO	60 AT 30	56 ~~ ~	·. ~	61	75		6	84		69	13		702
					··	×	-A G:	IG A	SA AC	ST C	נג גיו	W T	ra G	KA 27	C GA	C A	702 UA CGC
λз	n Tr	וא עד	A Hi	s T	r Se	r Al	a Va	al Az	g Se	r Le	u Ly	/# La	eu Va	 ea f	 n ls	 p Ly	A CGC
		71	1		72	0											
~-	- A1	T GT	T TC	C II	C CY	ТТА	C TA	C CA	y cc				C AC	A CA	, L CYC	ေထ	756 T GCC
Il	e Il	e Va	l Se	r Ph	e Hi	 s Ту:	 - Ty	r Gl	 u Pr	o Ph							T GCC
		76					_			- 1,1	u Dy	a rn	e Th	r Hi	3 Glr	Gl;	y Ala
GA	A TGG	GT	C	CC	77. 2 ATY	- ~	~	78: TO COM			79			801	L		810 TGG
								- GI	L AG	a GTT	r AA	TC	G AA'	r ccc	GAC	GAJ	TGG
Glu	Tr	Val	l Asr	Pro	) Ile	Pro	Pro	o Val	LArg	Val	Ly	Tr	P λει	i Gly	Glu	Gli	Trp
GAA	3.7~7	819		100	828	}		837	ı		846	5		855			864
			. CAA		- AGA	AGT	CAI	TIC	. YYY	TAC	GIC	AG:	GAC	TGG	GCA	λAG	864 CAA
Glu	Ile	λση	Gln	Ile	Arg	Ser	His	Phe	Lys	Tyr	Val	Sex	λs	JED 	Ala	Lys	Gln
		873			882			0.01									
AAT	AAC	GTA	CCA	ATC	TIT	CLL	GGT	. GYY	TTC	GGT			TCA	AAA	GCA	GAC	918 ATG
Asn	λsn	Val	Pro	Ile	Phe	Leu	Gly	Glu	Phe	Glv	λla	7V1		Lys		~	ATG
		927			936								261	rys	VIT	Asp	Het
GAC	TCA	AGG	GTT	λλG	TCC	100	GNA	945	~~~		954	_		963			972
								VQ1	GIG	AGA	AAA	ATG	GCG	963 GAA	GAA	TTT	GGA
Asp	Ser	Arg	Val	Lys	Trp	Thr	Glu	Ser	Val	λrg	Lvs	Mer		Glu			
		981			990									GIU	GTÜ	Pue	Gly
TTT	TCA	TAC	GCG	ፐልጥ	TCC	CAA	delen.	999						1017		- :	1026
								TGT	GCA	GGA	LLT	GGC	ATA	1017 TAC	Gat	λGA	TGG
Phe	Ser	Tyr	λla	Tyr	Trp	Glu	Phe	Суз	Ala	Gly	Phe	Gly	Ile	Tyr	Asn		~~~
	1	.035		1	044		,	1052		_							
TCT	СХА	AAC	TOG	ATC	GAA	CCA	327	CCR	101		1062	_		1071 ACA		1	1080
									~~~	oc I	010	GTT	GGC	ACX	CCC	λλλ	GλG
Ser	Gln	Asn	Trp	Ile	Glu	Pro	Leu	Ala	Thr	Ala	Val	Val	Gly	Thr	Gly	 Lys	Glu
AAT																	

Figure 13b(Continued)

Thermotoga maritima Pullulanase (6GP3)

5' ATG GAT CTT ACA	18	27	36	4 5 s
5' ATG GAT CTT ACA	ANG GIG GGG	ATC ATA GTG 1	AGG CTG AAC C	AG TGG CAG GCA AA
Met Asp Leu Thr	Lys Val Glv	Ile The Val		lu Trp Gln Ala Ly:
		are the val ;	ra ren vau d	lu Trp Gln Ala Ly:
63	72	81	90	99
CAC GIG CCA AAA	SAC AGG TTC)	ATA GAG ATA A	AA GAC GGA A	99 108 AG GCT GAA GTG TGG
Asp Val Ala Lys	SD Arg Phe I	'la Clu Tl	~ -	
-	-p ing life 1	re Cin lie L	As b Cla r	ys Ala Glu Val Trp
117	126	135	144	153
ATA CTC CAG GGA G	itg gaa gag a	TT TTC TAC G	AA AAA CCA CI	153 162 AC ACA TOT COO AGA
Ile Leu Gin Giv u				- TET TET CCC AGA
200 GH GIY V	ar Gra Gra I	le Phe Tyr G	lu Lys Pro As	p Thr Ser Pro Arg
171	180	100		
ATC TTC TTC GCA C	AG GCA AGG TO	G AAC AAG GT	באם האתר פאם פרי	207 216
Ile Pho Pho No Ale of			~	TITT CTG ACC AAT
Ile Phe Phe Ala G	in Ala Arg Se	er Asn Lys Va	l Ile Glu Al	a Phe Leu Thr Asn
225	234	244		
CCT CTG GAT ACG A	A AAG AAA GA	A CTC TTC AAG	252 3 GTT ACT CT	261 270
Drn 1/23 3			our act Gr	L CAC GGA YYY GYG
Pro Val Asp Thr Ly	'a Lya Lya G1	u Leu Phe Ly:	Val Thr Val	Asp Gly Lys Gly
279	288	207		
ATT CCC GTC TCA AG	A GTG GAA AA	297 G GCC GAT CCC	306	315 324
			ACG GAC AT	GAC GTG ACG AAC
Ile Pro Val Ser Ar	Val Glu Ly	Ala Asp Pro	Thr Asp Ile	Acr Val mb- a
333	342		-	TOP VAL THE ASH
TAC GTG AGA ATC GT	יאר מרות באר בי הרות האום	351	360	369 378
TAC GTG AGA ATC GT		·	GAN GAN GAC	CTC AGA AAA GAC
Tyr Val Arg Ile Val	l Leu Ser Glu	Ser Leu Lys	Glu Glu Asp	For her to
387				per vid PAR VED
GTG GAA CTG ATC ATI	396	405	414	423 432
GTG GAA CTG ATC ATI	INC	AAA CCG GCA	YCY CLC YLC	ATG ATG GAG ATC
Val Glu Leu Ile Ile	Glu Gly Tyr	Lys Pro Ala	Arg Val Tie	
			was agr 116	mer Met Glu Ile
441 CTG GAC GAC TAG TAG	450	459	468	477 485
CTG GAC GAC TAC TAT	INC GAT GGA	GAG CTC GGA	GCC GTA TAT	TCT CCA GAG AAG
Leu Asp Asp Tyr Tyr	TYT Asp Glv	Glu Leu Clu		
		sed GIA	ALA VAL TYT	Ser Pro Glu Lys
495	504	513	522	531 540
ACG ATA TTC AGA GTC	TGG TCC CCC	GTT TCT AAG	TCG GTA AAG	GIG CIT CIC TIC
Thr Ile Phe Arg Val	Tro Ser Pen	Val C *		
- 2 / 441	b par LLO	AUT SOL FAR	Trp Val Lys	Val Leu Leu Phe

Figure 14a

Thermotoga maritima Pullulanase (5GP3)	(continued)
549 550	
THE SON WALL ACK CALL COCK MAD CALC.	585 594
Lys Asn Gly Glu Asp Thr Glu Pro Tyr Gla Val	THE GAN TAC ANG GGA
Lys Asn Gly Glu Asp Thr Glu Pro Tyr Gln Val Val Asn	Met Glu Tyr Lys Gly
603 612	
AAC GGG GTC TGG GAA GCG GTT GTT GTA GGC GAT CTC GAC	GGA GTG TTC TAC CTC
Asn Gly Val Trp Glu Ala Val Val Glu Gly Asp Leu Asp	
657	GIY Val Phe Tyr Leu
TAT CAG CTG GAA AAC TAC GGA AAC ATG ACC	693 702
TAT CAG CTG GAA AAC TAC GGA AAG ATC AGA ACA ACC GTC	GAT CCT TAT TCG AAA
Tyr Gln Leu Glu Asn Tyr Gly Lys Ile Arg Thr Thr Val	LSD PIO TVI Ser Live
711 720 770	
GCG GTT TAC GCA AAC AAC CAA GAG AGC GCC GTT GTG AAT C	747 756 TT GCC AGG AGA AND
Ala Val Tyr Ala Asn Asn Gln Glu Gar Ala Wal	ACA AAC
Ala Val Tyr Ala Asn Asn Glm Glu Ser Ala Val Val Asn L	eu Ala Arg Thr Asn
765 774 783 792	801 810
CCA GAA GGA TGG GAA AAC GAC AGG GGA CCG AAA ATC GAA G	GA TAC GAA GAC GCG
Pro Glu Gly Trp Glu Asn Asp Arg Gly Pro Lys Ile Glu G	ly Tyr Glu Asp Ala
819 979	
ATA ATC TAT GAA ATA CAC ATA GCG GAC ATC ACA GGA CTC G	855 864
Ile Ile Tyr Glu Ile His Ile Ala Asp Ile Thr Gly Leu Gl	
	u Asn Ser Gly Val
873 882 891 900	909 918
AAA AAC AAA GGC CTC TAT CTC GGG CTC ACC GAA GAA AAC AC	G AAA GGA CCG GGC
Lys Asn Lys Gly Leu Tyr Leu Gly Leu Thr Glu Glu Asn Th	r Lys Gly Pro Gly
927 936 045	
GGT GTG ACA ACA GGC CTT TCG CAC CTT GTG GAA CTC GGT GT	963 972 T ACA CAC COM CAM
Gly Val Thr Thr Gly Len Sar Win I am	
Gly Val Thr Thr Gly Leu Ser His Leu Val Glu Leu Gly Va	l Thr His Val His
981 990 999 1008	1017 1026
ATA CTT CCT TTC TTT GAT TTC TAC ACA GCC GAC GAA CTC GA	T AAA GAT TTC GAG
Ile Leu Pro Phe Phe Asp Phe Tyr Thr Gly Asp Glu Leu Ax	The Age of the Commercial Commerc
1035 1044 1055	
ANG TAC TAC AND TEG GET TAC GAT COT TAC CTG TTC ATG GT	1071 1080
LYS TVT TVT Len Tom Clu D	- CCG GAG GGC AGA
Lys Tyr Tyr Asn Trp Cly Tyr Asp Pro Tyr Leu Phe Met Va.	l Pro Glu Gly Arg

Figure 14b(Continued)

Thermotoga maritima Pullulanase (6GP3) (continued)
TAC TCA ACC CAM CO. 1107 1116 1125
THE
Tyr Ser Thr Asp Pro Lys Asn Pro His Thr Arg Ile Arg Glu Val Lys Glu Met
1100
Val Lys Ala Leu His Lys His Gly Ile Gly Val Ile Man
The Net Val Phe Pro
CAC ACC TAC CCT NT 223
CAC ACC TAC GGT ATA GGC GAA CTC TCT GCG TTC GAT CAG ACG GTG CCG TAC TAC His Thr Tyr Gly Ile Gly Glu Leu Ser Ala Phe Arm Clar
His Thr Tyr Gly Ile Gly Glu Leu Ser Ala Phe Asp Gln Thr Val Pro Tyr Tyr
1451 1360 400
TTC TAC AGA ATC CAC AND ACC 1203 1278 1287 1287
Phe Tyr Arg Ile Asp Lys Thr Gly Ala Tyr Lou Are Gi
Phe Tyr Arg Ile Asp Lys Thr Gly Ala Tyr Leu Asn Glu Ser Gly Cys Gly Asn
115 ALC GCA AGC GA1 AGA COO 100 100 100 100 100 100 100 100 100 1
Val Ile Ala Ser Glu Arg Pro Met Met Arg Iva Dh.
Val Ile Ala Ser Glu Arg Pro Met Met Arg Lys Phe Ile Val Asp Thr Val Thr
1359 1360
TAC TGG GTA AAG GAG TAT CAC ATA GAC GGA TTC AGG TTC GAT CAG ATG GGT CTC
Tyr Trp Val Lys Glu Tyr His Ile Asp Gly Pho
Tyr Trp Val Lys Glu Tyr His Ile Asp Gly Phe Arg Phe Asp Gln Met Gly Leu
1413 4400
ATC GAC AAA AAG ACA AMO OMA
The Asp Lys Lys Thr Met Leu Glu Val Glu Asp Lys Lys Thr Met Lys Lys Thr Met Leu Glu Val Glu Asp Lys Lys Thr Met Lys Lys Thr Met Lys Thr Me
Ile Asp Lys Lys Thr Met Leu Glu Val Glu Arg Ala Leu His Lys Ile Asp Pro
1467 1476
ACT ATC ATT CTC TAC GGC GAA CCG TGG GGT GGA TGG GGA GCA CCG ATC AGG TTT
Thr Ile Ile Leu Tyr Gly Glu Pro Tro Gly Cly To Gly Cly
ory dry trp Gly Ala Pro Ile Arg Phe
157! 1536
SIY LYS Ser Asp Val Ala GIV Thr His Val Ala Si
Ala Ala Phe Ash Asp Glu Phe Arg
1575 1584
GAC GCA ATA AGG GGT TCC GTG TTC AAC CCG AGC GTC AAG GGA TTC GTC ATG GGA
ASP Ala Ile Arg Gly Ser Val Phe Ann Pro Ser Val
asp Ala Ile Arg Gly Ser Val Phe Asn Pro Ser Val Lys Gly Phe Val Met Gly

Figure 14C(Continued)

Thermotoga maritima Pullulanase (6GP3) (continued)

1629 163B 1647 GGA TAC GGA AAG GAA ACC AAG ATC AAA AGG GGT GTT GGT GGA AGC ATA AAC TAC Gly Tyr Gly Lys Glu Thr Lys Ile Lys Arg Gly Val Val Gly Ser Ile Asn Tyr 1683 1692 1701 GAC GGA AAA CTC ATC AAA AGT TTC GCC CTT GAT CCA GAA GAA ACT ATA AAC TAC --- --- --- --- --- --- --- --- --- --- --- --- --- --- ---Asp Gly Lys Leu Ile Lys Ser Phe Ala Leu Asp Pro Glu Glu Thr Ile Asn Tyr 1737 1746 1755 GCA GCG TGT CAC GAC AAC CAC ACA CTG TGG GAC AAG AAC TAC CTT GCC GCC AAA Ala Ala Cys His Asp Asn His Thr Leu Trp Asp Lys Asn Tyr Leu Ala Ala Lys 1791 1800 1809 GCT GAT ANG ANA ANG GAN TGG ACC GAN GAN GAN CTG ANA ANC GCC CAG ANA CTG --- --- --- --- --- --- --- --- --- --- --- --- --- --- --- ---Ala Asp Lys Lys Glu Trp Thr Glu Glu Leu Lys Asn Ala Gln Lys Leu 1845 1854 1863 COT GGT GCG ATA CTT CTC ACT TCT CAA GGT GTT CCT TTC CTC CAC GGA GGG CAG 1872 Ala Gly Ala Ile Leu Leu Thr Ser Gln Gly Val Pro Phe Leu His Gly Gly Gln 1899 1908 1917 GAC TTC TGC AGG ACG AAT TTC AAC GAC AAC TCC TAC AAC GCC CCT ATC TCG --- --- --- --- --- --- --- --- --- --- --- --- --- --- --- --- ---Asp Phe Cys Arg Thr Thr Asn Phe Asn Asp Asn Ser Tyr Asn Ala Pro Ile Ser 1953 1962 1971 ATA AAC GGC TTC GAT TAC GAA AGA AAA CTT CAG TTC ATA GAC GTG TTC AAT TAC 1980 Ile Asn Gly Phe Asp Tyr Glu Arg Lys Leu Gln Phe Ile Asp Val Phe Asn Tyr 2007 2016 2025 CAC AAG GGT CTC ATA AAA CTC AGA AAA GAA CAC CCT GCT TTC AGG CTG AAA AAC 2034 --- --- --- --- --- --- --- --- --- --- --- --- --- --- --- --- ---His Lys Gly Leu Ile Lys Leu Arg Lys Glu His Pro Ala Phe Arg Leu Lys Asn 2070 2079 GCT GAA GAG ATC AAA AAA CAC CTG GAA TTT CTC CCG GGC GGG AGA AGA ATA GTT --- --- --- --- --- --- --- --- --- --- ---Ala Glu Glu Ile Lys Lys His Leu Glu Phe Leu Pro Gly Gly Arg Arg Ile Val 2124 2133 GCG TTC ATG CTT AAA GAC CAC GCA GGT GGT GAT CCC TGG AAA GAC ATC GTG GTG --- --- --- --- --- --- --- --- --- --- --- --- --- --- --- --- ---Ala Phe Met Leu Lys Asp His Ala Gly Gly Asp Pro Trp Lys Asp Ile Val Val

Figure 14d(Continued)

Thermotoga maritima Fullulanase (6GP3) (continued)

2169 2178 2187 2196 2205 2214
ATT TAC AAT GGA AAC TTA GAG AAG ACA ACA TAC AAA CTG CCA GAA GGA AAA TGG
Ile Tyr Asn Gly Asn Leu Glu Lys Thr Thr Tyr Lys Leu Pro Glu Gly Lys Trp

2223 2232 2241 2250 2259 2268
AAT GTG GTT GTG AAC AGC CAG AAA CC GGA ACA GAA GTG ATA GAA ACC GTC GAA

AAT GTG GTT GTG AAC AGC CAG AAA CC GGA ACA GAA GTG ATA GAA ACC GTC GAA
Asn Val Val Val Asn Ser Gln Lys Ala Gly Thr Glu Val Ile Glu Thr Val Glu

GGA ACA ATA GAA CTC GAT CCG CTT TCC GCG TAC GTT CTG TAC AGA GAG TGA 3'
Gly Thr Ile Glu Leu Asp Pro Leu Ser Ale Tyr Val Leu Tyr Arg Glu ***

Figure 140(Continued)

Figure 15a Thermotoga maritima MSB8 (Clone # 6GP2) Glycosidase

1 CTT TTA TTG ATC GTT GAG CTC TCT TTC GTT CTC TTT GCA AGT GAC GAG TTC Leu Leu Leu Ile Val Glu Leu Ser Phe Val Leu Phe Ala Ser Asp Glu Phe

GTG AAA GTG GAA AAC GGA AAA TTC GCT CTG AAC GGA AAA GAA TTC AGA TTC Val Lys Val Glu Asn Gly Lys Phe Ala Leu Asn Gly Lys Glu Phe Arg Phe

ATT GGA AGC AAC AAC TAC TAC ATG CAC TAC AAG AGC AAC GGA ATG ATA GAC Ile Gly Ser Asn Asn Tyr Tyr Met His Tyr Lys Ser Asn Gly Met Ile Asp

AGT GTT CTG GAG AGT GCC AGA GAC ATG GGT ATA AAG GTC CTC AGA ATC TGG Ser Val Leu Glu Ser Ala Arg Asp Met Gly Ile Lys Val Leu Arg Ile Trp

GGT TTC CTC GAC GGG GAG AGT TAC TGC AGA GAC AAG AAC ACC TAC ATG CAT Gly Phe Leu Asp Gly Glu Ser Tyr Cys Arg Asp Lys Asn Thr Tyr Met His

CCT GAG CCC GGT GTT TTC GGG GTG CCA GAA GGA ATA TCG AAC GCC CAG AGC Pro Glu Pro Gly Val Pne Gly Val Pro Glu Gly Ile Ser Asn Ala Gln Ser

GGT TTC GAA AGA CTC GAC TAC ACA GTT GCG AAA GCG AAA GAA CTC GGT ATA Gly Phe Glu Arg Leu Asp Tyr Thr Val Ala Lys Ala Lys Glu Leu Gly Ile

AAA CTT GTC ATT GTT CTT GTG AAC AAC TGG GAC GAC TTC GGT GGA ATG AAC Lys Leu Val lle Val Leu Val Asn Asn Trp Asp Asp Phe Gly Gly Met Asn

CAG TAC GTG AGG TGG TTT GGA GGA ACC CAT CAC GAC GAT TTC TAC AGA GAT Gln Tyr Val Arg Trp Phe Gly Gly Thr His His Asp Asp Phe Tyr Arg Asp

GAG AAG ATC AAA GAA GAG TAC AAA AAG TAC GTC TCC TTT CTC GTA AAC CAT Glu Lys Ile Lys Glu Glu Tyr Lys Lys Tyr Val Ser Phe Leu Val Asn His

GTC AAT ACC TAC ACG GGA GTT CCT TAC AGG GAA GAG CCC ACC ATC ATG GCC Val Asn Thr Tyr Thr Gly Val Pro Tyr Arg Glu Glu Pro Thr Ile Met Ala

TGG GAG CTT GCA AAC GAA CCG CGC TGT GAG ACG GAC AAA TCG GGG AAC ACG TTP Glu Leu Ala Asn Glu Pro Arg Cys Glu Thr Asp Lys Ser Gly Asn Thr

CTC GTT GAG TGG GTG AAG GAG ATG AGC TCC TAC ATA AAG AGT CTG GAT CCC Leu Val Glu Trp Val Lys Glu Met Ser Ser Tyr Ile Lys Ser Leu Asp Pro

AAC CAC CTC GTG GCT GTG GGG GAC GAA GGA TTC TTC AGC AAC TAC GAA GGA Asn His Leu Val Ala Val Gly Asp Glu Gly Phe Phe Ser Asn Tyr Glu Gly

TTC AAA CCT TAC GGT GGA GAA GCC GAG TGG GCC TAC AAC GGC TGG TCC GGT Phe Lys Pro Tyr Gly Glu Ala Glu Trp Ala Tyr Asn Gly Trp Ser Gly

GTT GAC TGG AAG AAG CTC CTT TCG ATA GAG ACG GTG GAC TTC GGC ACG TTC Val Asp Trp Lys Lys Leu Leu Ser Ile Glu Thr Val Asp Phe Gly Thr Phe

CAC CTC TAT CCG TCC CAC TGG GGT GTC AGT CCA GAG AAC TAT GCC CAG TGG His Leu Tyr Pro Ser His Trp Gly Val Ser Pro Glu Asn Tyr Ala Gln Trp

GGA GCG AAG TGG ATA GAA GAC CAC ATA AAG ATC GCA AAA GAG ATC GGA AAA Gly Ala Lys Trp Ile Glu Asp His Ile Lys Ile Ala Lys Glu Ile Gly Lys

CCC GTT GTT CTG GAA GAA TAT GGA ATT CCA AAG AGT GCG CCA GTT AAC AGA Pro Val Val Leu Glu Glu Tyr Gly Ile Pro Lys Ser Ala Pro Val Asn Arg

ACG GCC ATC TAC AGA CTC TGG AAC GAT CTG GTC TAC GAT CTC GGT GGA GAT Thr Ala Ile Tyr Arg Leu Trp Asn Asp Leu Val Tyr Asp Leu Gly Gly Asp

GGA GCG ATG TTC TGG ATG CTC GCG GGA ATC GGG GAA GGT TCG GAC AGA GAC Gly Ala Met Phe Trp Met Leu Ala Gly Ile Gly Glu Gly Ser Asp Arg Asp

GAG AGA GGG TAC TAT CCG GAC TAC GAC GGT TTC AGA ATA GTG AAC GAC GAC Glu Arg Gly Tyr Tyr Pro Asp Tyr Asp Gly Phe Arg Ile Val Asn Asp Asp

AGT CCA GAA GCG GAA CTG ATA AGA GAA TAC GCG AAG CTG TTC AAC ACA GGT Ser Pro Glu Ala Glu Leu Ile Arg Glu Tyr Ala Lys Leu Phe Asn Thr Gly

GAA GAC ATA AGA GAA GAC ACC TGC TCT TTC ATC CTT CCA AAA GAC GGC ATG Glu Asp Ile Arg Glu Asp Thr Cys Ser Phe Ile Leu Pro Lys Asp Gly Met

GAG ATC AAA AAG ACC GTG GAA GTG AGG GCT GGT GTT TTC GAC TAC AGC AAC

Figure 15b (continued)

Glu Ile Lys Lys Thr Val Glu Val Arg Ala Gly Val Phe Asp Tyr Ser Asn

ACG TTT GAA AAG TTG TCT GTC AAA GTC GAA GAT CTG GTT TTT GAA AAT GAG Thr Phe Glu Lys Leu Ser Val Lys Val Glu Asp Leu Val Phe Glu Asn Glu

ATA GAG CAT CTC GGA TAC GGA ATT TAC GGC TTT GAT CTC GAC ACA ACC CGG Ile Glu His Leu Gly Tyr Gly Ile Tyr Gly Phe Asp Leu Asp Thr Thr Arg

ATC CCG GAT GGA GAA CAT GAA ATG TTC CTT GAA GGC CAC TTT CAG GGA AAA Ile Pro Asp Gly Glu His Glu Met Phe Leu Glu Gly His Phe Gln Gly Lys

ACG GTG AAA GAC TCT ATC AAA GCG AAA GTG GTG AAC GAA GCA CGG TAC GTG Thr Val Lys Asp Ser Ile Lys Ala Lys Val Val Asn Glu Ala Arg Tyr Val

CTC GCA GAG GAA GTT GAT TTT TCC TCT CCA GAA GAG GTG AAA AAC TGG TGG Leu Ala Glu Glu Val Asp Phe Ser Ser Pro Glu Glu Val Lys Asn Trp Trp

AAC AGC GGA ACC TGG CAG GCA GAG TTC GGG TCA CCT GAC ATT GAA TGG AAC Asn Ser Gly Thr Trp Gln Ala Glu Phe Gly Ser Pro Asp Ile Glu Trp Asn

GGT GAG GTG GGA AAT GGA GCA CTG CAG CTG AAC GTG AAA CTG CCC GGA AAG Gly Glu Val Gly Asn Gly Ala Leu Gln Leu Asn Val Lys Leu Pro Gly Lys

AGC GAC TGG GAA GAA GTG AGA GTA GCA AGG AAG TTC GAA AGA CTC TCA GAA Ser Asp Trp Glu Glu Val Arg Val Ala Arg Lys Phe Glu Arg Leu Ser Glu

TGT GAG ATC CTC GAG TAC GAC ATC TAC ATT CCA AAC GTC GAG GGA CTC AAG Cys Glu Ile Leu Glu Tyr Asp Ile Tyr Ile Pro Asn Val Glu Gly Leu Lys

GGA AGG TTG AGG CCG TAC GCG GTT CTG AAC CCC GGC TGG GTG AAG ATA GGC Gly Arg Leu Arg Pro Tyr Ala Val Leu Asn Pro Gly Trp Val Lys Ile Gly

CTC GAC ATG AAC AAC GCG AAC GTG GAA AGT GCG GAG ATC ATC ACT TTC GGC Leu Asp Met Asn Asn Ala Asn Val Glu Ser Ala Glu Ile Ile Thr Phe Gly

GGA AAA GAG TAC AGA AGA TTC CAT GTA AGA ATT GAG TTC GAC AGA ACA GCG Gly Lys Glu Tyr Arg Arg Phe His Val Arg Ile Glu Phe Asp Arg Thr Ala

Figure 15C(continued)

GGG GTG AAA GAA CTT CAC ATA GGA GTT GTC GGT GAT CAT CTG AGG TAC GAT Gly Val Lys Glu Leu His Ile Gly Val Val Gly Asp His Leu Arg Tyr Asp

GGA CCG ATT TTC ATC GAT AAT GTG AGA CTT TAT AAA AGA ACA GGA GGT ATG Gly Pro Ile Phe Ile Asp Asn Val Arg Leu Tyr Lys Arg Thr Gly Gly Met

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END

Figure 15d(continued)

Figure No. 16(Thermotoga maritima MSB8(6gb4)

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0.1	1 9	L GI	u Ar	g G1	u Ph	e Gl	u Ph	e Ly	s Gl	u As	p Va	l Ly	s Gl	u G	ly c	lu	Arg	Val	As	o Le	V	'a 1		
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81	Phe	Gli	1 Gl	y Va	l As	p Thi	r Le	ı Şe:	r Ası	o Vai	l Tv	r Lei	ו גם	n C1	31 G	-1	PAC	CTT	GG/	A AG	CA	CC	300	
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301	GAA	GAC	ATO	3 ጥጥ(ገ አጥ/	- GNG																		
101	Glu	Asr	Mer	Dhe	- ni	GAC	TA	CGC	TTC	GAT	GTC	ACC	3 AA	C GT	G T	TG A	AA	GAA	AAG	AA:	rc	AC	350	
				- F116	116	e Glu	Tyr	Arg	Phe	: Asp	Val	. Thi	: Ası	n Va	l L	eu I	ys	Glu	Lys	Ası	1 Ні	is	120	
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361	CTG	AAG	GTG	TAC	: ATA	AAA	TCT	CCC	ATC	AGA	GTT	, cca	AA.	A AC	T C1	rc g	AG (CAG	חממ	ም ክረ			420	
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481	GGA	TCC	CAC	maa																				
161	GGA Glv	7	SAC.	TGG	GGT	GCC	AGA	ATC	GTT	ACA	AGC	GGT	ATT	TGG	AA :	A C	CC C	TC	TAC	CTC	GA	G	540	
	Gly	rrþ	Asp	Trp	Gly	Ala	Arg	Ile	Val	Thr	Ser	Gly	Ile	Trp	Ly	s P	ro V	al '	Tyr	Leu	Gl	u	180	
541	GTG Val	TAC	AGG	GCA	CGT	CTT	CAG	GAT	TCA	ACG	GCT	TAT	CTG	TTG	G A	A C.	יידי ר	3 C (200			_		
181	Val	Tyr	Arg	Ala	Arg	Leu	Gln	Asp	Ser	Thr	Ala	Tvr	Leu	Leu	G1	n C.		AG (-3 -3	AAA	GA'	Г	600	
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601	GCC	CTT	GTG	AGG	GTG	ממ	CCT	TTC	cm.															
201	GCC Ala	Leu	Val	Ara	Val	Anc.	001	110	GIA	CAC	GGG	GAA	GGA	AAT	C.L.	C A	T G	TG (GAA	GTT	TA	r	660	
	Ala			7129	val	ASII	GIY	Phe	Vai	His	Gly	Glu	Gly	Asn	Le	u I]	e V	al (3 <u>1</u> u	Val	Ty	ŗ	220	
	GTA Val	AAC	GGT	GAA	AAG	ATA	GGG	GAG	TTT	CCT	GTT	CTT	GAA	AAG	AA:	C GC	A G	AA .	226	CTC	TT	_	720	
221	Val	Asn	Gly	Glu	Lys	Ile	Gly	Glu	Phe	Pro	Val	Leu	Glu	Lvs	Ası	n GI	v	3., 1		Lau	Db.	-		
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721	GAT Asp	GGA	GTG	TTC	CAC	CTG	AAA	GAT	GTG	an	CTA	TOO	m>~		_									
241	Asp	Gly	Val	Phe	His	Len	Live	A c.	Val	Lica	CIA	100	TAT	CCG	TG	G AJ	C C	TG (GGG	AAA	CCC	3	780	
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781 TAC CTG TAC GAT TTC GTT TTC CTG TTG AND AND THE	
781 TAC CTG TAC GAT TTC GTT TTC GTG TTG AAA GAC TTA AAC GGA GAG ATC TAC AGA GAA (261 Tyr Leu Tyr Asp Phe Val Phe Val Leu Lug Asp Tac Tac AGA GAA (GAA 840
261 Tyr Leu Tyr Asp Phe Val Phe Val Leu Lys Asp Leu Asn Gly Glu Ile Tyr Arg Glu (Glu 280
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841 AAG AAA ATC GGT TTG AGA AGA GTC AGA ATC GTT CAG GAG CCC GAT GAA GAA GGA AAA A	CT 900
281 Lys Lys Ile Gly Leu Arg Arg Val Arg Ile Val Gln Glu Pro Asp Glu Glu Gly Lys T	hr 300
THE GAM ALC AAC GGT GAG AAA GTC TTC COM ALC	CB 0.50
301 Phe Ile Phe Glu Ile Asn Gly Glu Lys Val Phe Ala Lys Gly Ala Asn Trp Ile Pro Se	
961 GAA AAC ATC CTC ACG TGG TTG AAG GAG GAA GAT TAC GAA AAG CTC GTC AAA ATG GCA AG	
321 Glu Asn Ile Leu Thr Trp Leu Lys Glu Glu Asp Tyr Glu Lys Leu Val Lys Met Ala Ar	G 1020
AND AND AND AND CITC AGG CITC TOO OF	
341 Ser Ala Asn Met Asn Met Leu Arg Val Trp Gly Gly Gly Ile Tyr Glu Arg Glu Ile Pho	C 1080
1081 TAC AGA CTC TGT GAT GAA CTC GGT ATC ATG GTG TGG CAG GAT TTC ATG TAC GCG TGT CTT	
361 Tyr Arg Leu Cys Asp Glu Leu Gly Ile Met Val Trp Gln Asp Phe Met Tyr Ala Cys Leu	1140
1141 GAA TAT CCG GAT CAT CTT CCG TGG TTC AGA AAA CTC GCG AAC GAA GAG GCA AGA AAG ATT	
381 Glu Tyr Pro Asp His Leu Pro Trp Phe Arg Lys Leu Ala Asn Glu Glu Ala Arg Lys Ile	
	400
1201 GTG AGA AAA CTC AGA TAC CAT CCC TCC ATT GTT CTC TGG TGC GGA AAC AAC GAA AAC AAC	
401 Val Arg Lys Leu Arg Tyr His Pro Ser Ile Val Leu Trp Cys Gly Asn Asn Glu Asn Asn	1260
	420
1261 TGG GGA TTC GAT GAA TGG GGA AAT ATG GCC AGA AAA GTG GAT GGT ATC AAC CTC GGA AAC	
421 Trp Gly Phe Asp Glu Trp Gly Asn Met Ala Arg Lys Val Asp Gly Ile Asn Leu Gly Asn	1320
	440
1321 AGG CTC TAC CTC TTC GAT TTT CCT GAG ATT TGT GCC GAA GAA GAC CCG TCC ACT CCC TAT	
441 Arg Leu Tyr Leu Phe Asp Phe Pro Glu Ile Cys Ala Glu Glu Asp Pro Ser Thr Pro Tyr	1380
	460
1381 TGG CCA TCC AGT CCA TAC GGC GGT GAA AAA GCG AAC AGC GAA AAG GAA GGA GAC AGG CAC 461 Trp Pro Ser Ser Pro Tyr Gly Gly Gly Loo Tyr	
461 Trp Pro Ser Ser Pro Tyr Gly Gly Glu Lys Ala Asn Ser Glu Lys Glu Gly Asp Arg His	1440
	480
1441 GTC TGG TAC GTG TGG AGT GGC TGG ATG AAC TAC GAA AAC TAC GAA AAA GAC ACC GGA AGG	
481 Val Trp Tyr Val Trp Ser Gly Trp Met Asn Tyr Glu Asn Tyr Glu Lys Asp Thr Gly Arg	1500
And Tyr Glu Lys Asp Thr Gly Arg	500
1501 TTC ATC AGC GAG TTT GGA TTT CAG GGT GCT CCC CAT CCA GAG ACG ATA GAG TTC TTT TCA	
501 Phe Ile Ser Glu Phe Gly Phe Gln Gly Ala Pro His Pro Glu Thr Ile Glu Phe Phe Ser	1560
The Fig. als Pro Glu Thr Ile Glu Phe Phe Ser	520
1561 AAA CCC GAG GAA AGA GAG ATA TTC CAT GGG GTG	
1561 AAA CCC GAG GAA AGA GAG ATA TTC CAT CCC GTC ATG CTG AAG CAC AAC AAA CAG GTG GAA	1620
o and the his proval Met Leu Lys His Asn Lys Gln Val Glu	540
Figure 16b(continued)	

162 54		GA.	CAG	GAA	A A G	A TT	G AT	C AG	G TT	C AT.	A TT	C GG	A A	AT '	III	GGA	A A A	TO	אמ ד	۸ ۵				
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																								=
168	1 A	GT :	TT	GTG	TA	CT	G TCC	CAC	CTC	: AAC	CAC		a er											
56	1 Se	er I	he	Val.	Тут	Let	ı Ser	Gln	Leu	Asn	Glr	1 Al:	9 (C)		1 -	ATC	AAG	TTC	GG	I G	TT (GAA	CAC	1740
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1741	L TG	GC	GA	AGC	AGG	AAG	TAC	444	ACC.	ccc	000													
581	Tr	pΑ	rg	Ser	Arq	Lvs	TAC	Lva	The	11-	GGC	GCI	· CT	CT	TC '	TGG	CAG	TTC	AAC	GA	C A	GC	TGG	1800
					-	•	туг	٠, ٠	****	n_a	GIY	Ala	Le	u Pl	ne '	Trp	Gln	Phe	Asn	As	p S	er	Trp	600
1801	CC	3 G:	rc '	TTC	AGC	TGG	TCC	CC.																
601	Pro	V V	al 1	Phe	Ser	Trn	TCC	BIR	GTC	GAT	TAC	TTC	AAA	A AC	G	CCC	AAA	GCT	CTC	TA	C T	AC '	TAT	1860
							Ser	A14	Vai	Asp	Tyr	Phe	Lys	Ar	g F	ro	ГЛа	Ala	Leu	Ty	r T	yr '	Tyr	620
1861	GCG	AG	A A	(C) -	P-P-C	TOTAL C	:											•						
621	Ala	Ar	σ A	. במים)he	Dha	GCT	GAA	GTT	CTA	CCC	GTT	TTG	AA	G A	AG /	AGA (GAC	AAÇ	AAA	LA A	'A C	AA	1920
			J -		116	FILE	Ala	GIU	Val 1	Leu	Pro	Val	Leu	Ly	s L	ys A	Arg /	Asp .	Asn	Lys	: Il	e 0	lu	640
1921																								
641	T.eu	T.m.		IG G	iGT (GAG	CGA :	rcr (GAG (GA (GAC 2	AAA	AGA	AGT	r c	TC T	CI (AG (GCT	TGC	AG	c c	TA	1980
	200	пе	. V.	al G	TÀ (Glu.	Arg s	Ser (3lu c	ly ,	Asp 1	Lys	Arg	Ser	Le	eu S	er o	iln /	la	Cys	Se	r L	eu	660
1981																								****
661	CGA	GAA	L GA	AA G	GG ,	AGA A	AAA C	GT A	TT C	GA A	AA C	SAC :	ΓTΑ	CAG	AA	C G	GT A	CT C	מר :	ACC.	NG:		~~	2040
901	Arg	Glu	G)	lu G	ly A	Arg I	∵ya G	ly I	le A	rg L	ув А	sp 1	Leu	Gln	As	n G	ly T	hr F	ro !	Ser	Arc	- C(2040
																	•	_				, A.	9	680
2041	TGT						205	5																
681	Сув	Glu	Ph	e GI	ly E	nđ	685																	

Figure 16 c(continued)

Figure No. 12 Bankia gouldi (37gp4)

	1 A	TG A	LAA A	AAA A	LAP (TA C	TA A	TG I	TT A	AA A	GG (TT A	ACG 1	י דמי	- A.T.					TG CTG	
	1 M	et I	ys I	Lys)	Asn I	eu L	eu M	et P	he I	vs A	ro I	.e 7	'h= 7	· · · ·	-14	CCT :	rrg 1	TTT 1	TA A	NTG CTG let Leu	60
										,,,,,,	-9 .	,	414 1	yr I	eu i	Pro I	Leu F	he L	eu M	let Leu	20
6	1 C	יי יי	C 2 C	·~ .	~ ~																
2:	·			-1A A	GT T	CA G	TA G	CT C	AA T	CT C	CT G	TA G	AA A	AA C	AT C	GC C	GT T	TA C	AA G	TT GAC	120
-		eu s	er L	eu s	er S	er V	al A	La G	ln S	er P	ro V	al G	lu L	γs H	is G	ly A	rg L	eu G	ln V	TT GAC	40
																					10
121	. G G	A A	AC C	GC A	TT C	IT AJ	AT GO	G TO	T G	SA GA	AA A	IT A	G A	יר די	TB C	CT C	-m .			C TTT	
41	Gl	y A	sn A	rg I	le L	eu A≤	n Al	a Se	r G	.v G1	u I	le Ti	17 S4	- T		1- 0	JI A	AC AC	C CI	C TTT	180
										•					eu A	Id G.	LY AS	in Se	r Le	u Phe	60
181	TG	G AG	T A	AT G	ساب (5)	a ca	c .c														
61	Tr	ri Se	γ Ac	וא מי		L	- AC	C TC	C GA	T TT	T TA	T AA	T GC	A GA	A A	T GI	T GA	T TT	T TT	A GCA	240
		,		··· ^	La GI	у АБ	p Th	r Se	r As	p Ph	e Ty	r As	n Al	a Gl	u Th	ır Va	l As	p Ph	e Le	A GCA u Ala	80
241	GA	A AA	C TG	G AA	T AG	C TC	A CT	r at	r ag	A AT	A GC	T AT	G GG	C GT	а аа	A GA	A AA	r ra	GA'	r GGC	200
81	Glı	ı As	n Tr	p As	n Se	r Se	r Lei	ı Ile	a Ar	3 Il	a Al	a Me	t Gl	y Va	l Lv	s G1:	ıı Aqı	Tr	A DA	r GGC o Gly	300
																	- 115		vol	, GIA	100
301	GGA	AA.	r gg	C TA	T AT	I GAT	AGI	cca	: כאר	CAC	. Chi									GAT	
101	Gly	Ası	1 Gl	y Tv	r Ile	a Asr	Ser	Dro	Gle	Clu	01-	· OA	4 GC	AA,	A AT	T AG	A AAA	GTT	ATI	GAT	360
				• •					, 611	GIL	L GII	1 GIL	1 Ala	Lys	3 Il	e Arç	Lys	Val	Ile	Asp	120
361	GCN	CO																			
121	NIA	. 601	AT.	r GC:	I AAC	GGC	ATA	TAT	GTA	ATA	ATA .	GAC	TGG	CAC	: AC	CAC	GAA	GCA	GAG	TTA	420
121	νīτα	ATS	1116	e Ala	a Asr	Gly	Ile	Tyr	Val	Ile	Ile	Asp	Trp	His	Thi	His	Glu	Ala	Glu	Leu	140
421	TAC	ACA	GAT	GAC	GCT	GTT	GAC	TTT	TTT	ACC	AGA	ATG	GCA	GAC	מידים י	י מידי	CCA	C2.00			
141	Tyr	Thr	Asp	Glu	Ala	Val	Asp	Phe	Phe	Thr	Arq	Met	Ala	Agn	LAN	7	Clas	GAT	ACT	CCC	480
									-		-				שבט	TYL	GIY	Asp	Inr	Pro	160
481	AAT	GTA	ATG	דבד ;		ስ ጥጥ	TAT	220	a												
161	Asn	Val	Met	ጥν	GAA	T10	T	AAC	GAG	ccr	ATA	TAC	CAA	AGT	TGG	CCT	GTT	ATT	AAG	AAT	540
				-,-	Glu	116	Tyr	ASI	GIU	Pro	Ile	Tyr	Gln	Ser	Trp	Pro	Val	Ile	Lys	Asn	180
E 4 1																					
541	TAT	GCA	GAG	CAA	GTA	ATT	GCT	GGT	ATA	CGT	TCT	AAA	GAC	CCA	GAT	AAT	TTA	АТА	ATT	GTA	600
181	Tyr	ALa	Glu	Gln	Val	Ile	Ala	Gly	Ile	Arg	Ser	Lys	Asp	Pro	Asp	Asn	Leu	Ile	Ile	Va1	200
															_						100
601	GGT	ACT	AGC	AAT	TAT	TCT	CAG	CAA	GTT	GAT	GTA	GCA	ጥ ር እ	CCZ	010						
201	Gly	Thr	Ser	Asn	Tyr	Ser	Gln	Gln	Val	Agn	Val	Ala	Cam	GCA	GAC	CCA	ATA	TCT	GAT	ACT	660
					-					лэр	•41	wra	ser	ATA	Asp	Pro	Ile	Ser	Asp	Thr	220
661	AAT	GTC	CCN	Th				_													
221	Asn	Ual	Bl.	TAT	ACT	TTA	CAT	TTT	TAT	GCA	GCA	TTT	AAC	CCG	CAT	GAT	AAC	TTA	AGA	AAT	720
-	- 1011	• a I	wrg	t.Az	Thr	Leu	His	Phe	Tyr	Ala	Ala	Phe	Asn	Pro	His	Asp	Asn	Leu	Arg	Asn	240
721	GTA	GCA	CAG	ACA	GCA	TTA	GAT	AAT	AAT	GTT	GCT	TTG	TTT	GTT	ACN	ממם	TCC	CC=		3 mm	200
241	Val	Ala	Gln	Thr	Ala	Leu	Asp	Asn	naA	Val	Ala	Leu	Pha	Val	The	GAA	166	GGT	ACA	ATT	780
							-				•	Leu	FIIC	VAI	ınr	Glu	Trp	Gly	Thr	Ile	260

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781 TTA AAT ACC GGA CAA GGA GAA CCA GAC AAA GAA AGC ACT AAT ACT TGG ATG GCC TTT TT	
261 Leu Asn Thr Gly Gln Gly Glu Pro Asp Lys Glu Ser Thr Asn Thr Trp Met Ala Phe Leu	G 840
Die Ser inr Asn Thr Trp Met Ala Phe Let	u 280
841 AAA GAA AAA GGT ATA AGT CAC GCT AAT TCC TOT TO	
841 AAA GAA AAA GGT ATA AGT CAC GCT AAT TGG TCT TTG AGT GAC AAA GCT TTT CCT GAA ACA 281 Lys Glu Lys Gly Ile Ser His Ala Asn Trp Ser Leu Ser Asp Lys Ala Phe Pro Glu Thr	900
The Ser Leu Ser Asp Lys Ala Phe Pro Glu Thr	300
901 GGG TCT GTA GTT CAA GCA GGA CAA GCT GTA TOT ON	
901 GGG TCT GTA GTT CAA GCA GGA CAA GGT GTA TOT GGT TTA ATT AGC AAT AAA CTT ACA GCC 301 Gly Ser Val Val Gln Ala Gly Gln Gly Val Ser Gly Leu Ile Ser Asn Lys Leu Thr Ala	960
, ser Gly Leu Ile Ser Asn Lys Leu Thr Ala	320
961 TCT GGT GAA ATT GTA AAA AAC ATC ATC CAA AAC ATC	
961 TCT GGT GAA ATT GTA AAA AAC ATC ATC CAA AAC TGG GAT ACA GAG ACC TCT ACA GGA CCT 321 Ser Gly Glu Ile Val Lys Asn Ile Ile Gln Asn Trp Asp Thr Glu Thr Ser Thr Gly Pro	1020
of Ash Trp Asp Thr Glu Thr Ser Thr Gly Pro	340
1021 AAA ACA ACA CAA TGT AGT ACT ATA GAA TCT ATT AGT	
1021 AAA ACA ACA CAA TGT AGT ACT ATA GAA TGT ATT AGA GCT GCA ATG GAA ACA GCA CAA GCA 341 Lys Thr Thr Gln Cys Ser Thr Ile Glu Cys Ile Arg Ala Ala Met Glu Thr Ala Gln Ala	1080
tot of the Arg Ala Met Glu Thr Ala Gln Ala	360
1081 GGA GAT GAA ATT ATA ATT GCC CCT GGA AAC TAC AND	
1081 GGA GAT GAA ATT ATA ATT GCC CCT GGA AAC TAC AAT TTT CAA GAC AAG ATA CAA GGT GCC 361 Gly Asp Glu Ile Ile Ile Ala Pro Gly Asn Tyr Asn Phe Gln Asp Lys Ile Gln Gly Ala	1140
Ash Phe Gin Asp Lys Ile Gln Gly Ala	380
1141 TTT AAC CGT AGT GTT TAC CTT TAT GGT AGT GCT AAC GGA AAC AGT ACA AAC CCT ATT ATA 381 Phe Asn Arg Ser Val Tyr Leu Tyr Gly Ser Ala Acc GGA AAC AGT ACA AAC CCT ATT ATA	
381 Phe Asn Arg Ser Val Tyr Leu Tyr Gly Ser Ala Asn Gly Asn Ser Thr Asn Pro Ile Ile	1200
	400
1201 TTA AGA GGC GAA AGC GCT ACA AAC CCT CCT GTT TTC TCA GGA TTA GAT TAT AAC AAT GGC	
401 Leu Arg Gly Glu Ser Ala Thr Asn Pro Pro Val Phe Ser Gly Leu Asp Tyr Asn Asn Gly	1260
	420
1261 TAC CTA TTA AGT ATT GAA GGT GAT TAT TGG AAT ATT AAA GAT ATA GAG TTT AAA ACT GGG	
421 Tyr Leu Leu Ser Ile Glu Gly Asp Tyr Trp Asn Ile Lys Asp Ile Glu Phe Lys Thr Gly	1320
	440
1321 TCT AAA GGT ATT GTT CTT GAC AAT TCT AAT GGT AGT AAA TTA AAA AAC CTT GTT GTT CAT :	
der Lys Leu Lys Ash Lys Ash Leu Lys Ash Ly	1380
	460
THE GOA GAA GAA GCT ATT CAC THE	
and all Asp Gly Ser Ser Asp Asp Com The	1440
1441 TCC Non-new	480
1441 TGC ACT ATA TAC AAT ACA GGT AGA ACT AAA CCT GGT TTT GGT GAA GGT TTA TAT GTA GGC 1	
The GIV GIV Con The services	.500
	500
THE GAL AAA GGA CAA CAT GAC ACT TAT CAN AGE	r.c.
And Cys Ash Ash Ash The The	560 520
	\$20
ACC GIT GGA CCC AAT GTA ACA CCC COT COT	
of val Asp val Lvs Glu Glu man	620
try fine Met Asn	540

Figure 17b(continued)

1	621 ACT ATT ATA AGA AAT TGC GTG TTT TCT GCA GAA GGA ATT TCA GGA GAA AAT AGC TCA GAT 541 Thr lie lie Arg Asn Cys Val Dhe Ser No. Cl. Cl. Co.	
:	541 Thr Ile Ile Arg Asn Cys Val Phe Ser No Chy Ci	1680
	541 Thr Ile Ile Arg Asn Cys Val Phe Ser Ala Glu Gly Ile Ser Gly Glu Asn Ser Ser Asp	560
16		
	ALL GALLIA AAA GGA GCC TAT CCT TTT CTA TAG	
	661 Ala Phe Ile Asp Leu Lys Gly Ala Tyr Gly Phe Val Tyr Arg Asn Thr Phe Asn Val Asp	1740
	and the Phe Asn Val Asp	580
17	41 GGT TCT GAA GTA ATA AAT ACT CCA CTA CAA	
5	41 GGT TCT GAA GTA ATA AAT ACT GGA GTA GAC TTT TTA GAT AGA GGT ACA GGA TTT AAT ACA	1800
	81 Gly Ser Glu Val Ile Asn Thr Gly Val Asp Phe Leu Asp Arg Gly Thr Gly Phe Asn Thr	
		600
180	ALL GOA ATA TIT GAA AAT ACA TAT AAC COM	
60	Ol Gly Phe Arg Asn Ala Ile Phe Glu Asn Thr Tyr Asn Leu Gly Ser Arg Ala Ser Glu Ile	1860
	John Ded Gly Ser Arg Ala Ser Glu Ile	620
186	1 TCA ACT GCT CGT ARA ARA CAR COM TOTAL	
62	1 TCA ACT GCT CGT AAA AAA CAA GGT TCT CCT GAA CAA ACT CAC GTT TGG GAT AAT ATT AGA	1920
	1 Ser Thr Ala Arg Lys Lys Gln Gly Ser Pro Glu Gln Thr His Val Trp Asp Asn Ile Arg	640
100	·	
192	ALL ICI GIT GAT TIT CCA ATA ACT CAT CCT PC	
641	Asn Pro Asn Ser Val Asp Phe Pro Ile Ser Asp Gly Thr Glu Asn Leu Val Asn Lys Phe	1980
	The state of the s	660
1981	TGC CCA GAT TGG AAT ATA GAA CCA TGT AND GOT	
661	TGC CCA GAT TGG AAT ATA GAA CCA TGT AAT CCT GTA GAC GAA ACC AAC CAA GCA CCT ACA	2040
	. Cys Pro Asp Trp Asn Ile Glu Pro Cys Asn Pro Val Asp Glu Thr Asn Gln Ala Pro Thr	680
2041		
	THE CIR ICI CCT GTT AAC AAT ATT ACT TTA CTT	
901	ASH ASH ILE THE Leu Val Gly Tyr has to	100
		700
2101	GAA GTT AAT GCT ACT GAT GCA GAT GGA ACT ATT GAT AAT GTA AAA CTT TAT ATA GAT AAC 2.	
701	Glu Val Asn Ala Thr Asp Ala Asp Gly Thr Ile Asp Asn Val Lys Leu Tyr Ile Asp Asn	160
	and the Asp Ash Val Lys Leu Tyr Ile Asp Ash	720
2161	AAT TITL COM THE COM	
721	AAT TTA GTT AGG CAA ATA AAT TCT ACT TCA TAT AAA TGG GGC CAT TCT GAT TCT CCA AAT 22	220
	The Ash Ser Thr Ser Tyr Lys Trp Gly Wie Co-	
		740
2221	ACA GAT GAA CTT AAT GGT CTT ACA GAA GGA ACT TAT ACC TTA AAA GCA ATT GCA ACT GAT 22	
741	Thr Asp Glu Leu Asn Gly Leu Thr Glu Gly Thr Tyr Thr Leu Lys Ala Ile Ala Thr Asp	80
	The Leu Lys Ala He Ala Thr Asp	760
2281	AAC GAC GGG GGT TGT AGA TA	
761	AAC GAC GGG GCT TCT ACA GAA ACG CAA TTT ACG TTA ACT GTA ATA ACA GAA CAA AGT CCG 23	40
	off the Gin Phe Thr Leu Thr Val The The Classic	80
		~ •
2341	TCT GAG AAT TGT GAC TTT AAT ACA CCT TCT TCA ACT GGT TTA GAA GAT TTT GAC ATT AAA 24	
781	Ser Glu Asn Cys Asp Phe Asn Thr Pro Ser Ser Thr Gly Leu Glu Asp Phe Asp Ile Lys 8	00
	8 and only Led Glu Asp Phe Asp Ile Lys	00
2401	AAG TIT TCT AAC GIT TIT CAG TON	
	AAG TIT TCT AAC GIT TIT GAG TTA GGA TCT GGC GGA CCA TCT TTA AGT AAT TTA AAA ACA 24	60
	Figure 17C(continued)	

Figure 174(continued)

80	1 Lys Phe Ser									
246	L TIT ACT ATT	AAT TGG A	AT TCG	Cha mad				•		
82.				.,,-	U. y	ned Tyr	Gin Phe	Ser Ile	A AAC ACA AAC Asn Thr Asn	2520 840
252] 841	AAC GGT GTA (CCT GAT TA	AT TAT A	ITA AAT le Asn	TTA AAA	CCA AAA . Pro Lys :	ATT ACC	TTT CAG	TTT AAA AAT Phe Lys Asn	2580 860
2581 861	GCA AAT CCA G Ala Asn Pro G	AA ATA TC lu Ile Se	T ATT AG	GC AAT A er Asn S	AGC TTA A	TT CCT A	AT TTT	GAT GGT Asp Gly	GAT TAC TGG Asp Tyr Trp	2640 880
2641 881		AT AAC GG1	ייים יוממי	T 000 -						2700 900
2701	TTT AGT AAT GA Phe Ser Asn As	C GCT ACT	GCT CC	سن سنوس ا	· · · · · · · · · · · · · · · · · · ·	_				2760 920
2761 921	ATT ACT GAT GAT	T TCT AGT	ATT AAT	TTT AA	G CTT TAG s Leu Tŷi	CCT AN	r CCT GC	T TTA G à Leu A	AC GAA ACT	2820 940
2821 .	ATT TTT GTG AGC	GCT GAA	GAT GAA	AAA CTY	\ CC# ###				70	

Figure 17d(continued)

Figure No. 18 Pyrococcus furiosus VC1(7EG1)

13111 No. 140 Pyrococcus furiosus VC1 (7EG1)
leader sequence: amino acids 1-24
9
2/ 36
5' ATG AGC AAG AAA AAG TTC GTC ATC GTA TCT ATC TTA ACA ATC CTT TTA GTA CAG
Met Ser Lys Lys Lys Phe Val Ile Val Ser Ile Leu Thr Ile Leu Leu Val Gln
<u>.</u>
72 81 80 80
GCA ATA TAT TTT GTA GAA AAG TAT CAT ACC TCT GAG GAC AAG TCA ACT TCA AAT
Ala Ile Tyr Phe Val Glu Lys Tyr His Thr Ser Glu Asp Lys Ser Thr Ser Asn
117 126 135 144 153 162
ACC TCA TCT ACA CCA CCC CAA ACA ACA CTT TCC ACT ACC AAG GTT CTC AAG ATT
Thr Ser Ser Thr Pro Pro Gln Thr Thr Leu Ser Thr Thr Lys Val Leu Lys Ile
171 180 189 198 207 216
AGA TAC CCT GAT GAC GGT GAG TGG CCA GGA GCT CCT ATT GAT AAG GAT GGT GAT
Arg Tyr Pro Asp Asp Gly Glu Trp Pro Gly Ala Pro Ile Asp Lys Asp Gly Asp
225 234 243 252 261 270
GGG AAC CCA GAA TTC TAC ATT GAA ATA AAC CTA TGG AAC ATT CTT AAT GCT ACT
Gly Asn Pro Glu Phe Tyr Ile Glu Ile Asn Leu Trp Asn Ile Leu Asn Ala Thr
9
279 288 297 306 315 324
GGA TTT GCT GAG ATG ACG TAC AAT TTA ACC AGC GGC GTC CTT CAC TAC GTC CAA
Gly Phe Ala Glu Met Thr Tyr Asn Leu Thr Ser Gly Val Leu His Tyr Val Gln
333 342 351 360 369 378
CAA CTT GAC AAC ATT GTC TTG AGG GAT AGA AGT AAT TGG GTG CAT GGA TAC CCC
Gln Leu Asp Asn Ile Val Leu Arg Asp Arg Ser Asn Trp Val His Gly Tyr Pro
387 396 405 414 423 432
GAA ATA TTC TAT GGA AAC AAG CCA TGG AAT GCA AAC TAC GCA ACT GAT GGC CCA
Glu Ile Phe Tyr Gly Asn Lys Pro Trp Asn Ala Asn Tyr Ala Thr Asp Gly Pro
441 450 459 468 477 486
ATA CCA TTA CCC AGT AAA GTT TCA AAC CTA ACA GAC TTC TAT CTA ACA ATC
Ile Pro Leu Pro Ser Lys Val Ser Asn Leu Thr Asp Phe Tyr Leu The Ile Ser

Ile Pro Leu Pro Ser Lys Val Ser Asn Leu Thr Asp Phe Tyr Leu Thr Ile Ser

TAT AAA CTT GAG CCC AAG AAC GGC CTG CCA ATT AAC TTC GCA ATA GAA TCC TGG Tyr Lys Leu Glu Pro Lys Asn Gly Leu Pro Ile Asn Phe Ala Ile Glu Ser Trp

603 612 621 630 639 648 ATG ATA TAG TAT GAC GGA TTA CAA CCG GCT GGC TCC AAA GTT AAG GAG Met Ile Trp Ile Tyr Asp Gly Leu Gln Pro Ala Gly Ser Lys Val Lys Glu

ATT GTA GTC CCA ATA ATA GTT AAC GGA ACA CCA GTA AAT GCT ACA TTT GAA GTA ILe Val Val Pro Ile Ile Val Asn Gly Thr Pro Val Asn Ala Thr Phe Glu Val

THE FIGURE 152 STATE ST

765 774 783 792 801 810 AAA GAG GGA ACA GTG ACA ATT CCA TAC GGA GCA TTT ATA AGT GTT GCA GCC AAC Lys Glu Gly Thr Val Thr Ile Pro Tyr Gly Ala Phe Ile Ser Val Ala Ala Asn

 $873 \hspace{0.2cm} 882 \hspace{0.2cm} 891 \hspace{0.2cm} 900 \hspace{0.2cm} 518$ $ACT \hspace{0.2cm} GAG \hspace{0.2cm} TTT \hspace{0.2cm} GGA \hspace{0.2cm} ACG \hspace{0.2cm} CCAC \hspace{0.2cm} ACG \hspace{0.2cm} CCAC \hspace{0.2cm} CTAC \hspace{0.2cm} GAG \hspace{0.2cm} TGG \hspace{0.2cm} ATC \hspace{0.2cm} ACAC \hspace{$

AAC ATA ACA CTA ACT CCT CTA GAT AGA CCT CTT ATT TCC TAA 3'
Asn Ile Thr Leu Thr Pro Leu Asp Arg Pro Leu Ile Ser *

Figure 18b(continued)

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/22623

IPC(6) US CL	IPC(6) :C07H 21/04; C12N 1/20, 1/14, 5/00, 9/38, 9/42; C08B 30/04 US CL :435/207, 209, 252.3, 254.11, 274, 275, 320.1, 325; 536/23.2 According to International Patent Classification (IPC) or to both national classification and IPC										
	LDS SEARCHED										
Minimum o	Minimum documentation searched (classification system followed by classification symbols)										
U.S.: 435/207, 209, 252.3, 254.11, 274, 275, 320.1, 325; 536/23.2											
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched											
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)											
Please See Extra Sheet.											
:											
C. DOCUMENTS CONSIDERED TO BE RELEVANT											
Category*	Citation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.								
X GRABNITZ et al. Structure of the β-Glucosidase Gene bglA of Clostridium thermocellum: Sequence Analysis Reveals a Superfamily of Cellulases and β-Glycosidases Including Human Lactase/Phlorizin Hydrolase. Eur. J. Biochem. September 1991, Vol. 200, No. 2, pages 301-309, see entire document.											
X A	X VOORHORST et al. Characterization of the celB Gene Coding for β-Glucosidase from the Hyperthermophilic Archaeon Pyrococcus species I and III										
Furt	her documents are listed in the continuation of Box C.	See patent family annex.									
	or o	*T* later document published after the inter date and not in conflict with the appli	cation but cited to understand								
	becoment defining the general state of the art which is not considered be of particular relevance	the principle or theory underlying the	1								
	urlier document published on or efter the international filing data	"X" document of particular relevance; the considered novel or cannot be consider when the document is taken alone	red to involve an inventive step								
oit	ted to establish the publication date of another citation or other	"Y" document of particular relevance; the	claimed invention cannot be								
special reason (as specified) *O* document referring to an oral disclosure, use, exhibition or other means *O* document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art											
P document published prior to the international filing date but later than *&* document member of the same patent family the priority date claimed											
	Date of the actual completion of the international search Date of mailing of the international search report										
26 MARCH 1998 <u>2</u> 1 APR 1998											
Name and mailing address of the ISA/US Authorized officer											
Commissioner of Patents and Trademarks Box PCT LISA J. HOBBS. PH.D.											
_	Washington, D.C. 20231 Facsimile No. (703) 305-3230 Telephone No. (703) 308-0196										

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/22623

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	
1.	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2.	Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inte	emational Searching Authority found multiple inventions in this international application, as follows:
Please See Extra Sheet.	
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. X	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: 1-11, species I-III
4. 🔲	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark	on Protest The additional search fees were accompanied by the applicant's protest.
	No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/22623

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS and STN (Bioscience and Patent Indexes): Desulfurococc##, Staphylotherm##, Thermatoga, galactosidase#, glucosidase#, beta galactosidase#, beta glucosidase#. Genbank, EMBL, ESTs1-4, STS, N-Geneseq: Seq. ID Nos.: 1-3 and A-Geneseq, PIR, Swissprot: Seq ID Nos.: 15-17.

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This applies on contains claims directed to more than one species of the generic invention. These species are deemed to lack Unity of Invention because they are not so linked as to form a single inventive concept under PCT Rule 13.1. The species are as follows: there are 18 distinct enzymes disclosed in the description, as enumerated in Figs. 1-18 and Table 1.

The claims are deemed to correspond to the species listed above in the following manner: while all the claims form one Group for examination, each of the claims is generic to the 18 enzyme species disclosed.

The species listed above do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, the species lack the same or corresponding special technical features for the following reasons: each enzyme is a different product, thus has the special technical feature of the recited enzyme, which the other species lack.